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Murakami et al.

(54) METHOD OF PRODUCING **ELECTROPHOTOGRAPHIC** PHOTOSENSITIVE MEMBER, AND **EMULSION FOR A CHARGE** TRANSPORTING LAYER

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See application file for complete search history.

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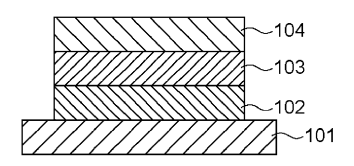
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(57)**ABSTRACT**

A method of producing an electrophotographic photosensitive member includes: preparing a solution including a charge transporting substance, and at least one compound selected from the group consisting of a fluorine-atomcontaining polyacrylate, a fluorine-atom-containing polymethacrylate, a polycarbonate having a siloxane bond, a polyester having a siloxane bond, a polystyrene having a siloxane bond, a silicone oil, a polyolefin, an aliphatic acid, an aliphatic acid amide and an aliphatic acid ester; preparing an emulsion by using the solution and water; forming a coat of the emulsion on a support; and heating the coat to form a charge transporting layer.

13 Claims, 2 Drawing Sheets



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FIG. 1A

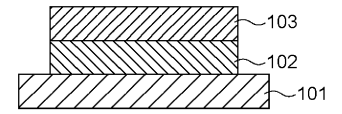


FIG. 1B

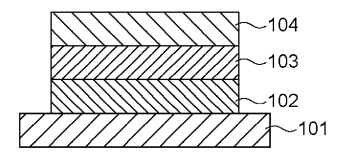
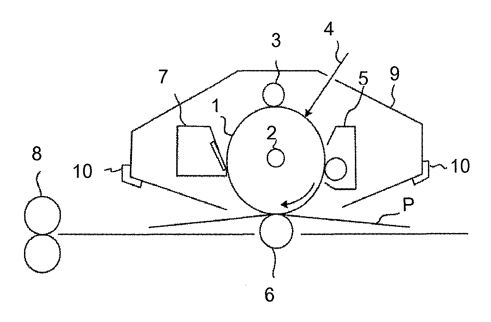


FIG. 2



METHOD OF PRODUCING ELECTROPHOTOGRAPHIC PHOTOSENSITIVE MEMBER, AND EMULSION FOR A CHARGE TRANSPORTING LAYER

TECHNICAL FIELD

The present invention relates to a method of producing an electrophotographic photosensitive member, and an emul- 10 sion for a charge transporting layer.

BACKGROUND ART

Electrophotographic photosensitive members to be mounted on electrophotographic apparatuses include organic electrophotographic photosensitive members containing an organic photoconductive substance (hereinafter, also referred to as an "electrophotographic photosensitive member"). The organic electrophotographic photosensitive members are currently a mainstream as an electrophotographic photosensitive member used in a process cartridge for the electrophotographic apparatus or the electrophotographic apparatus, and produced in a large scale. Among these electrophotographic photosensitive members, a laminate type electrophotographic photosensitive member is often used, of which properties are improved by separately providing the functions necessary for the electrophotographic photosensitive member in individual layers.

A method of producing the laminate type electrophotographic photosensitive member is usually used in which a functional material is dissolved in an organic solvent to prepare an application solution (coating solution), and the coating solution is applied onto a support. Among the layers in the laminate type electrophotographic photosensitive member, a charge transporting layer often demands durability. For this reason, the charge transporting layer has a film thickness of a coat relatively thicker than those of other layers. Accordingly, a large amount of the coating solution is used for the charge transporting layer, resulting in a large amount of the organic solvent to be used. In order to reduce the amount of the organic solvent to be used in production of the electrophotographic photosensitive member, the amount of the organic solvent to be used for the coating solution for a charge transporting layer is desirably reduced. To prepare the coating solution for a charge transporting layer, however, a halogen solvent or an aromatic organic solvent needs to be used because a charge transporting substance and a binder resin are highly soluble in the halogen solvent or the aromatic organic solvent. For this reason, the amount of the organic solvent to be used is difficult to reduce.

PTL 1 discloses an attempt to reduce a volatile substance and the amount of an organic solvent to be used in a coating solution for forming a charge transporting layer (coating solution for a charge transporting layer). PTL 1 discloses preparation of an emulsion type coating solution (emulsion) by forming an organic solution into oil droplets in water in which the organic solution is prepared by dissolving a substance included in a charge transporting layer in an organic solvent.

CITATION LIST

Patent Literature

PTL 1: Japanese Patent Application Laid-Open No. 2011-128213

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SUMMARY OF INVENTION

Technical Problem

As a result of research by the present inventors, however, it was found out that in the method of producing an electrophotographic photosensitive member disclosed in PTL 1 in which the emulsion is prepared, the emulsion is uniformly emulsified immediately after the preparation of the emulsion, but the liquid properties of the emulsion are reduced after the emulsion is left as it is for a long time.

The reason for this is thought as follows: the organic solution prepared by dissolving the substance included in a charge transporting layer in the organic solvent coalesces in water as the time has passes; this coalescence makes it difficult to form a stable state of oil droplets, leading to aggregation or sediment. Then, further improvement is desired from the viewpoint of reducing the amount of the organic solvent to be used and ensuring the stability of the coating solution for a charge transporting layer at the same time.

An object of the present invention is to provide a method of producing an electrophotographic photosensitive member in which the amount of an organic solvent to be used for a coating solution for a charge transporting layer is reduced, and the stability of the coating solution for a charge transporting layer after preservation for a long time is improved, enabling formation of a charge transporting layer having high uniformity.

Another object of the present invention is to provide a coating solution for a charge transporting layer having high stability after preservation for a long time.

Solution to Problem

The objects above are attained by the present invention below.

The present invention is a method of producing an electrophotographic photosensitive member which includes a support, and a charge transporting layer formed thereon, the method including: preparing a solution including: a charge transporting substance; and at least one compound selected from the group consisting of a fluorine-atom-containing polyacrylate, a fluorine-atom-containing polymethacrylate, a polycarbonate having a siloxane bond, a polyester having a siloxane bond, a polystyrene having a siloxane bond, a silicone oil, a polyolefin, an aliphatic acid, an aliphatic acid amide and an aliphatic acid ester; dispersing the solution in water to prepare an emulsion; forming a coat for the charge transporting layer by using the emulsion; and heating the coat to form the charge transporting layer.

Moreover, the present invention relates to an emulsion for a charge transporting layer in which a solution is dispersed in water, wherein the solution includes: a charge transporting substance; and at least one compound selected from the group consisting of a fluorine-atom-containing polyacrylate, a fluorine-atom-containing polymethacrylate, a polycarbonate having a siloxane bond, a polyester having a siloxane bond, a polystyrene having a siloxane bond, a silicone oil, a polyolefin, an aliphatic acid, an aliphatic acid amide and an aliphatic acid ester.

Advantageous Effects of Invention

The present invention can provide a method of producing an electrophotographic photosensitive member in which the stability of the coating solution for a charge transporting

layer (emulsion) after preservation for a long time can be improved, enabling formation of a charge transporting layer having high uniformity. Moreover, the present invention can provide a coating solution for a charge transporting layer (emulsion) having high stability after preservation for a long 5 time.

Further features of the present invention will become apparent from the following description of exemplary embodiments with reference to the attached drawings.

Further features of the present invention will become ¹⁰ apparent from the following description of exemplary embodiments with reference to the attached drawings.

BRIEF DESCRIPTION OF DRAWINGS

FIGS. 1A and 1B are drawings showing an example of a layer configuration in an electrophotographic photosensitive member according to the present invention.

FIG. 2 is a drawing showing an example of a schematic configuration of an electrophotographic apparatus including ²⁰ a process cartridge having the electrophotographic photosensitive member according to the present invention.

DESCRIPTION OF EMBODIMENTS

As described above, the method of producing an electrophotographic photosensitive member according to the present invention includes: preparing a solution including: a charge transporting substance; and at least one compound selected from the group consisting of a fluorine-atom-containing polyacrylate, a fluorine-atom-containing polymethacrylate, a polycarbonate having a siloxane bond, a polyester having a siloxane bond, a polystyrene having a siloxane bond, a silicone oil, a polyolefin, an aliphatic acid, an aliphatic acid amide and an aliphatic acid ester; dispersing the solution in water to prepare an emulsion; forming a coat for the charge transporting layer by using the emulsion; and heating the coat to form the charge transporting layer.

The present inventors think the reason why the method of producing an electrophotographic photosensitive member 40 according to the present invention can improve the stability of the emulsion (coating solution for a charge transporting layer) after preservation for a long time, enabling formation of a charge transporting layer having high uniformity as follows.

In the present invention, in preparation of the solution containing the charge transporting substance, a solution further containing a compound that provides an effect of reducing surface energy (fluorine-atom-containing polyacrylate, fluorine-atom-containing polymethacrylate, polycarbonate having a siloxane bond, polyester having a siloxane bond, polystyrene having a siloxane bond, silicone oil, polyolefin, aliphatic acid, aliphatic acid amide, aliphatic acid ester) is prepared. By preparing an emulsion including the solution and water, the emulsion never aggregates (co-slesses) even if the emulsion is preserved for a long time. It is thought that this provides the effect of the present invention.

As the techniques described in PTL 1, a period for which the dispersion state of the emulsion is kept can be extended 60 by containing a large amount of a surfactant, but the oil droplet state (emulsion) may be difficult to keep. Then, it is thought that in the present invention, by addition of the compound that provides an effect of reducing surface energy above, the surface energy of the oil droplets in the emulsion 65 is reduced to reduce an aggregation (coalescence) force of the oil droplets, and thereby, aggregation (coalescence) of

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the oil droplets is suppressed. For this reason, aggregation of the emulsion is suppressed even after the emulsion is preserved for a long time, and stability of the emulsion is enhanced. Moreover, because aggregation of the emulsion caused by preservation for a long time is suppressed, use of even the emulsion after preservation for a long time allows formation of a charge transporting layer having high uniformity.

Hereinafter, the materials that form the electrophotographic photosensitive member produced by the production method above will be described.

The electrophotographic photosensitive member produced by the production method above is an electrophotographic photosensitive member including a support, and a charge transporting layer formed thereon. The electrophotographic photosensitive member can be a laminate type (function separate type) photosensitive layer in which a charge generating layer containing a charge generating substance and a charge transporting layer containing a charge transporting substance are separately provided. The laminate type photosensitive layer may be a normal layer type photosensitive layer in which the charge generating layer and the charge transporting layer are laminated in this order from the side of the support, or may be an inverted layer type photosensitive layer in which the charge transporting layer and the charge generating layer are laminated in this order from the side of the support. From the viewpoint of electrophotographic properties, the normal layer type photosensitive layer can be used.

FIGS. 1A and 1B are drawings showing an example of a layer configuration of the electrophotographic photosensitive member according to the present invention. In FIGS. 1A and 1B, a support 101, a charge generating layer 102, a charge transporting layer 103, and a protective layer 104 (second charge transporting layer) are shown. When necessary, an undercoat layer may be provided between the support 101 and the charge generating layer 102.

The charge transporting substance is a substance having a hole transporting ability. Examples of the charge transporting substance include triarylamine compounds or hydrazone compounds. Among these, use of the triarylamine compounds can be used from the viewpoint of improving the electrophotographic properties.

The specific examples of the charge transporting substance are shown below:

$$H_3C$$
 CH_3 H_3C CH_3

25

30

35

CTM-5

CTM-6

55

60

CTM-7

-continued

CTM-2 H_3C

CTM-3 15 C_6H_{13}

 C_6H_{13}

CTM-4

-continued

CH₃ 10 CTM-8

 H_3 H_3C CTM-9

The charge transporting substance may be used alone or 40 in combination.

As a material that forms the charge transporting layer, a binder resin may be contained.

Examples of the binder resin used for the charge trans-45 porting layer include styrene resins, acrylic resins, polycarbonate resins and polyester resins. Among these, polycarbonate resins or polyester resins can be used. Further, polycarbonate resins having a repeating structural unit represented by the following formula (B1) or polyester resins 50 having a repeating structural unit represented by the following formula (B2) can be used.

where R^{51} to R^{54} each independently represent a hydrogen atom or a methyl group; X3 represents a single bond, a 65 methylene group, an ethylidene group, a propylidene group, a phenylethylidene group, a cyclohexylidene group or an oxygen atom.

-continued

(B1-4)

(B2-1)

(B2-2)

where R^{55} to R^{58} each independently represent a hydrogen atom or a methyl group; X^4 represents a single bond, a methylene group, an ethylidene group, a propylidene group, a cyclohexylidene group or an oxygen atom; Y^3 represents an m-phenylene group or a p-phenylene group or a divalent group having two p-phenylene groups bonded with an oxygen atom.

Specific examples of the repeating structural unit represented by the formula (B1) are shown below:

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\$$

$$\begin{array}{c|c} & CH_3 & O & \\ \hline \\ C & CH_3 & O & C \\ \hline \\ CH_3 & O & C \\ \end{array}$$

$$\begin{array}{c|c} & & & & \\ & & & \\ O & & & \\ \end{array} \begin{array}{c} & & & \\ & & \\ \end{array} \begin{array}{c} & & \\ & \\ \end{array} \begin{array}{c} & & \\ & \\ \end{array} \begin{array}{c} & & \\ \end{array} \begin{array}{c} & & \\ \end{array} \begin{array}{c} & & \\ & \\ \end{array} \begin{array}{c} & &$$

$$-\begin{bmatrix} 0 & & & & & \\ & & & & & \\ 0 & & & & & \\ \end{bmatrix} - \begin{bmatrix} 0 & & & & \\ & & & & \\ \end{bmatrix} - \begin{bmatrix} 0 & & & & \\ & & & & \\ \end{bmatrix}$$

Specific examples of the repeating structural unit represented by the formula (B2) are shown below:

$$\begin{bmatrix} H_3C & CH_3 & CH_3 & O & C\\ CH_3 & CH_3 & CH_3 & O & C\\ CH_3 & CH_3 & CH_3 & O & C\\ CH_3 & CH_3 & CH_3 & C\\ CH_3 & CH_$$

-continued

(1)

These polycarbonate resins and polyester resins can be used alone, or can be used in combination by mixing or as a copolymer. The form of the copolymerization may be any form of block copolymerization, random copolymerization and alternating copolymerization. The polycarbonate resins ³⁵ and polyester resins above can have no siloxane bond because the effect of the present invention is obtained stably.

The weight average molecular weight of the binder resin is a weight average molecular weight in terms of polystyrene measured according to the standard method, specifically according to the method described in Japanese Patent Application Laid-Open No. 2007-079555.

In the present invention, examples of the fluorine-atom-containing polyacrylate and the fluorine-atom-containing bolymethacrylate include a compound having a repeating structural unit represented by the following formula (1):

$$\begin{bmatrix}
 & H \\
 & C \\
 & C$$

$$\begin{array}{c|c}
 & R^{11} \\
\hline
 & C \\
 & C \\$$

where R^{11} represents hydrogen or a methyl group; R^{12} represents an alkylene group, and can be an alkylene group having 1 to 4 carbon atoms; R^{13} represents a perfluoroalkyl group having 4 to 6 carbon atoms.

Hereinafter, specific examples of the repeating structural unit represented by the formula (1) are shown:

50 (1-3)
$$\begin{bmatrix}
 & & & H \\
 & & & C \\
 & & & C \\
 & & & & C
\end{bmatrix}$$

$$\begin{bmatrix}
 & & & H \\
 & & & C \\
 & & & C
\end{bmatrix}$$

$$\begin{bmatrix}
 & & & H \\
 & & & C
\end{bmatrix}$$

$$\begin{bmatrix}
 & & & H \\
 & & & C
\end{bmatrix}$$

$$\begin{bmatrix}
 & & & H \\
 & & & C
\end{bmatrix}$$

$$\begin{bmatrix}
 & & & C
\end{bmatrix}$$

$$\begin{bmatrix}
 & CH_3 \\
 & C \\$$

$$\begin{bmatrix}
 & CH_3 \\
 & C \\
 & C \\
 & CH_2
\end{bmatrix}$$

$$\begin{bmatrix}
 & CH_3 \\
 & C \\
 & CH_2
\end{bmatrix}$$

$$\begin{bmatrix}
 & CH_3 \\
 & C \\
 & CH_2
\end{bmatrix}$$

$$\begin{bmatrix}
 & CH_3 \\
 & C \\
 & CH_2
\end{bmatrix}$$

$$\begin{bmatrix}
 & CH_3 \\
 & C \\
 & CH_2
\end{bmatrix}$$

$$\begin{bmatrix}
 & CH_3 \\
 & C \\
 & CH_2
\end{bmatrix}$$

$$\begin{bmatrix}
 & CH_3 \\
 & C \\
 & CH_2
\end{bmatrix}$$

$$\begin{bmatrix}
 & CH_3 \\
 & C \\
 & CH_2
\end{bmatrix}$$

$$\begin{bmatrix}
 & CH_3 \\
 & C \\
 & CH_2
\end{bmatrix}$$

$$\begin{bmatrix}
 & CH_3 \\
 & C \\
 & CH_2
\end{bmatrix}$$

$$\begin{bmatrix}
 & CH_3 \\
 & CH_3
\end{bmatrix}$$

$$\begin{bmatrix}
 & CH_3 \\
 & C$$

$$\begin{bmatrix} & & & & & & & \\ & & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & \\ & & & & & \\ & & & \\ & & & & \\ & & & & \\ & & & \\ & & & & \\ & & & & \\ & & \\ & &$$

$$\begin{bmatrix} & & & & & & & & & \\ & & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\$$

$$\begin{bmatrix} & & & & & & H & & & \\ & & & & & & C & - CH_2 & & \\ & & & & & & C & - CH_2 & & \\ & & & & & & & C & - CH_2 &$$

$$\begin{bmatrix} \text{CH}_3 \\ \text{C} \\ \text{CC} \\ \text{C} \end{bmatrix} \\ \begin{bmatrix} \text{CH}_2 \\ \text{C} \\ \text{CCH}_2 \end{bmatrix} \\ \end{bmatrix}$$

The fluorine-atom-containing polyacrylates and fluorine-atom-containing polymethacrylates can be used alone, or can be used in combination by mixing or a copolymer. The form of copolymerization may be any form of block copolymerization, random copolymerization and alternating copolymerization.

In the emulsion according to the present invention, the content of the fluorine-atom-containing polyacrylate and the fluorine-atom-containing polymethacrylate can be not less than 0.1% by mass and not more than 1% by mass based on the total mass of the charge transporting substance and the binder resin. At a content within this range, the effect of stabilizing the emulsion by use of the fluorine-atom-containing polyacrylate and the fluorine-atom-containing polymethacrylate can be sufficiently obtained, and the effect of sufficient electrophotographic properties can be obtained.

Examples of the polycarbonate having a siloxane bond include polycarbonate A having a repeating structural unit represented by the following formula (2-1) and a repeating structural unit represented by the following formula (2-3), or polycarbonate B having a repeating structural unit represented by the following formula (2-2) and repeating structural unit represented by the following formula (2-3):

-continued

$$X^{1}$$
 X^{1}
 X^{32}
 X^{1}
 X^{32}
 X^{32}
 X^{32}
 X^{32}
 X^{33}
 X^{33}

In the formula (2-1), R^{14} to R^{17} each independently ²⁵ represent a methyl group or a phenyl group; m^1 represents the number of repetition of the structure enclosed in brackets, and the average of m^1 in the polycarbonate A ranges from 20 to 100. Further, the number of repetition of the structure enclosed in brackets m^1 is preferably within the range of $\pm 10\%$ of the value indicated by the average of the number of repetition of m^1 because the effect of the present invention is obtained stably.

In the formula (2-2), R^{18} to R^{29} each independently 35 represent a methyl group or a phenyl group; m^2 , m^3 , m^4 , and m^5 each independently represent the number of repetition of the structure enclosed in brackets, and the average of $m^2+m^3+m^4+m^5$ in the polycarbonate B ranges from 0 to 450; Z^1 and Z^2 each independently represent an ethylene group or a propylene group; Z^3 represents a single bond, an oxygen atom, an ethylene group or a propylene group. Further, the sum of the numbers of repetition of the structure enclosed in brackets $m^2+m^3+m^4+m^5$ is preferably within the range of $\pm 10\%$ of the value indicated by the average of the 45 number of repetition of $m^2+m^3+m^4+m^5$ because the effect of the present invention is obtained stably.

In the formula (2-3), X¹ represents a single bond, a methylene group, an ethylidene group, a propylidene group, a phenylethylidene group, a cyclohexylidene group or an 50 oxygen atom; R³⁰ to R³³ each independently represent a hydrogen atom or a methyl group.

Hereinafter, specific examples of the repeating structural unit represented by the formula (2-1) are shown. In Table 1, the average of m^1 represents the average of m^1 in the 55 polycarbonate A.

TABLE 1

			_		
Repeating structural unit represented by formula (2-1)	R ¹⁴	R ¹⁵	R ¹⁶	R ¹⁷	Average of m ¹
Repeating structural unit example (2-1-1)	Methyl group	Methyl group	Methyl group	Methyl group	20

TABLE 1-continued

(2-3)

30	Repeating structural unit represented by formula (2-1)	R ¹⁴	R^{15}	R ¹⁶	R ¹⁷	Average of m ¹
	Repeating structural	Methyl group	Methyl group	Methyl group	Methyl group	40
35	unit example (2-1-2)	Strap	Second	Secup	Seed	
	Repeating	Methyl	Methyl	Methyl	Methyl	60
	structural	group	group	group	group	
40	unit example					
	(2-1-3)					
	Repeating	Methyl	Methyl	Methyl	Methyl	100
	structural	group	group	group	group	
45	unit example					
	(2-1-4)					
	Repeating	Methyl	Methyl	Phenyl	Methyl	40
	structural	group	group	group	group	
50	unit example					
	(2-1-5)					
	Repeating	Phenyl	Methyl	Methyl	Methyl	40
	structural	group	group	group	group	
55	unit example					
	(2-1-6)					
	Repeating	Phenyl	Methyl	Phenyl	Methyl	40
	structural	group	group	group	group	
60	unit example					
00	(2-1-7)					

Hereinafter, specific examples of the repeating structural unit represented by the formula (2-2) are shown. In Table 2, the sum of m², m³, m⁴, and m⁵ represents the average of m²+m³+m⁴+m⁵ in the polycarbonate B.

TABLE 2

Repeating structural unit represented by formula (2-2)	R	¹⁸ -R ²⁹	Z^1	Z^2	Z^3	m ²	m ³	m ⁴	m ⁵
Repeating structural unit	R ²⁰ ,R ²⁷ -R ²⁹	9: Methyl group	Propylene	Propylene	Ethylene group	0	0	0	0
example (2-2-1) Repeating structural unit example (2-2-2)	R ¹⁸ -R ²⁹ :	Methyl group	group Propylene group	group Propylene group	Ethylene group	1	1	1	100
Repeating structural unit example (2-2-3)	R^{18} - R^{29} :	Methyl group	Ethylene group	Ethylene group	Ethylene group	1	1	1	200
Repeating structural unit example (2-2-4)	R ¹⁸ -R ²⁹ :	Methyl group	Propylene group	Propylene group	Ethylene group	1	1	1	400
Repeating structural unit example (2-2-5)	R ¹⁸ -R ²⁹ :	Methyl group	Propylene	Propylene group	Ethylene group	20	20	20	20
Repeating structural unit example (2-2-6)	R^{18} - R^{29} :	Methyl group	Ethylene group	Ethylene group	Ethylene group	100	100	50	200
Repeating structural unit example (2-2-7)	R ¹⁸ -R ²⁹ :	Methyl group	Propylene group	Propylene group	Ethylene group	150	150	50	100
Repeating structural unit example (2-2-8)	R ²⁵ ,R ²⁷ -R ²⁹ :	R ¹⁸ ,R ²¹ ,R ²⁴ ,R ²⁶ : Phenylene	Propylene group	Propylene group	Ethylene group	20	20	20	20
Repeating structural unit example (2-2-9)	Methyl group R ²⁰ ,R ²⁷ -R ²⁹ : Methyl group	group R ²⁵ ,R ²⁶ : Phenylene group	Propylene group	Propylene group	Ethylene group	0	0	0	100
Repeating structural unit example (2-2-10)	R ¹⁸ -R ²⁴ ,R ²⁷ -I	R ²⁹ : Methyl group	Propylene	Propylene	Single bond	20	20	100	0
Repeating structural unit example (2-2-11)	R ¹⁸ -R ²⁴ ,R ²⁷ -I	R ²⁹ : Methyl group	group Propylene group	group Propylene group	Single bond	100	100	100	0
Repeating structural unit example (2-2-12)	R ¹⁸ -R ²⁹ :	Methyl group	Propylene group	Propylene group	Propylene group	100	100	100	100
Repeating structural unit	R ¹⁸ -R ²⁹ :	Methyl group	Propylene	Propylene	Propylene	20	20	20	20
example (2-2-13) Repeating structural unit example (2-2-14)	R ¹⁸ -R ²⁴ ,R ²⁷ -I	R ²⁹ : Methyl group	group Ethylene group	group Ethylene group	group Single bond	20	20	100	0
Repeating structural unit	R ¹⁸ -R ²⁴ ,R ²⁷ -I	R ²⁹ : Methyl group	Ethylene	Ethylene group	Single bond	150	150	150	0
example (2-2-15) Repeating structural unit example (2-2-16)	R ¹⁸ -R ²⁹ :	Methyl group	group Ethylene group	Ethylene group	Ethylene group	20	20	20	20

55

Specific examples of the repeating structural unit represented by the formula (2-3) include the repeating structural units represented by the formulas (B1-1) to (B1-8). The present invention is not limited to these.

In the polycarbonate having a siloxane bond, the poly-40 carbonate A and the polycarbonate B can have a terminal structure represented by the following formula (2-4) in one terminal or both terminals. In the case where the polycarbonate A and the polycarbonate B have the terminal struc- 45 etition enclosed in brackets; the average of m11 in the ture represented by the formula (2-4) in one terminal, a molecular weight adjuster (terminal terminator) is used to terminate the other terminal. Examples of the molecular weight adjuster include phenol, para-cumylphenol, paratert-butylphenol, and benzoic acid. Among these, phenol and para-tert-butylphenol can be used. In this case, the other terminal structure is a terminal structure represented by the following formula (2-5) or the following formula (2-6):

O (2-4)
$$(CH_{2})_{3} \leftarrow \begin{pmatrix} R^{61} \\ I \\ Si - O \\ I \\ R^{63} \end{pmatrix}_{m^{11}} \begin{pmatrix} CH_{3} \\ Si - CH_{3} \\ I \\ CH_{3} \end{pmatrix}$$

$$(2-5)^{-65}$$

-continued

$$\begin{array}{c} CH_3 \\ C \\ C \\ CH_3 \\ CH_3 \end{array}$$

In the formula (2-4), m11 represents the number of reppolycarbonate A or the polycarbonate B ranges from 20 to 100; R⁶¹ and R⁶² each independently represent a methyl group or a phenyl group.

Hereinafter, specific examples of the terminal structure ⁵⁰ represented by the formula (2-4) are shown:

$$\begin{array}{c|c} & & & & & & & & \\ & & & & & & & \\ & & & & & & \\ & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & &$$

-continued

The polycarbonates having a siloxane bond can be used alone, or can be used in combination by mixing.

The content of the polycarbonate having a siloxane bond in the emulsion can be not less than 0.1% by mass and not more than 5% by mass based on the total mass of the charge 20 transporting substance and the binder resin. At a content within this range, the effect of stability of the emulsion by used of the polycarbonate having a siloxane bond can be sufficiently obtained, and the effect of sufficient electrophotographic properties can be obtained.

Examples of the polyester having a siloxane bond include polyester C having a repeating structural unit represented by the following formula (3-1) and a repeating structural unit represented by the following formula (3-2):

-continued

In the formula (3-1), R³⁴ to R³⁷ each independently represent a methyl group or a phenyl group; Y¹ represents a meta-phenylene group, a para-phenylene group, or a bivalent group having two para-phenylene groups bonded with an oxygen atom; m⁶ represents the number of repetition of the structure enclosed in brackets, and the average of m⁶ in the polyester C ranges from 20 to 100.

In the formula (3-2), R^{38} to R^{41} each independently represent a hydrogen atom or a methyl group; X^2 represents a single bond, a methylene group, an ethylidene group, a propylidene group, a cyclohexylidene group or an oxygen atom; Y^2 represents a meta-phenylene group, a para-phenylene group or a bivalent group having two para-phenylene groups bonded with an oxygen atom.

Hereinafter, specific examples of the repeating structural unit represented by the formula (3-1) are shown. In Table 3, the average of m^6 represents the average of m^6 in the polyester C.

TABLE 3

Repeating structural unit represented by formula (3-1)	R ³⁴	R ³⁵	R ³⁶	R ³⁷	Average of m ⁶	Y ¹
Repeating structural unit example (3-1-1)	Methyl group	Methyl group	Methyl group	Methyl group	20	p-Phenylene group
Repeating structural unit example (3-1-2)	Methyl group	Methyl group	Methyl group	Methyl group	40	p-Phenylene group
Repeating structural unit example (3-1-3)	Methyl group	Methyl group	Methyl group	Methyl group	60	p-Phenylene group
Repeating structural unit example (3-1-4)	Methyl group	Methyl group	Methyl group	Methyl group	100	p-Phenylene group
Repeating structural unit example (3-1-5)	Methyl group	Methyl group	Phenyl group	Methyl group	40	p-Phenylene group
Repeating structural unit example (3-1-6)	Phenyl group	Methyl group	Methyl group	Methyl group	40	p-Phenylene group
Repeating structural unit example (3-1-7)	Phenyl group	Methyl group	Phenyl group	Methyl group	40	p-Phenylene group
Repeating structural unit example (3-1-8)	Methyl group	Methyl group	Methyl group	Methyl group	20	m-Phenylene group
Repeating structural unit example (3-1-9)	Methyl group	Methyl group	Methyl group	Methyl group	40	m-Phenylene group
Repeating structural unit example (3-1-10)	Methyl group	Methyl group	Methyl group	Methyl group	60	m-Phenylene group
Repeating structural unit example (3-1-11)	Methyl group	Methyl group	Methyl group	Methyl group	100	m-Phenylene group
Repeating structural unit example (3-1-12)	Methyl group	Methyl group	Phenyl group	Methyl group	40	m-Phenylene group

TABLE 3-continued

Repeating structural unit epresented by formula (3-1)	R^{34}	R^{35}	R ³⁶	R^{37}	Average of m ⁶	Y^1
Repeating structural unit example (3-1-13)	Phenyl group	Methyl group	Methyl group	Methyl group	40	m-Phenylene group
Repeating structural unit example (3-1-14)	Phenyl group	Methyl group	Phenyl group	Methyl group	40	m-Phenylene group
Repeating structural unit example (3-1-15)	Methyl group	Methyl group	Methyl group	Methyl group	20	
Repeating structural unit example (3-1-16)	Methyl group	Methyl group	Methyl group	Methyl group	40	-0-
Repeating structural unit example (3-1-17)	Methyl group	Methyl group	Methyl group	Methyl group	60	
Repeating structural unit example (3-1-18)	Methyl group	Methyl group	Methyl group	Methyl group	100	
Repeating structural unit example (3-1-19)	Methyl group	Methyl group	Phenyl group	Methyl group	40	
Repeating structural unit example (3-1-20)	Phenyl group	Methyl group	Methyl group	Methyl group	40	
Repeating structural unit example (3-1-21)	Phenyl group	Methyl group	Phenyl group	Methyl group	40	

(3-4) 65

Specific examples of the repeating structural unit represented by the formula (3-2) include repeating structural units 40 represented by the formulas (B2-1) to (B2-6).

In the polyester having a siloxane bond, the polyester C may have a terminal structure represented by the formula (3-3) in one terminal or both terminals. In the case where the polyester C has the terminal structure represented by the 45 formula (3-3) in one terminal, a molecular weight adjuster (terminal terminator) is used to terminate the other terminal. Examples of the molecular weight adjuster include phenol, para-cumylphenol, para-tert-butylphenol, and benzoic acid. Among these, phenol and para-tert-butylphenol can be used. In this case, the other terminal structure is a terminal structure represented by the following formula (3-5) or the following formula (3-6):

$$(3-3)$$

$$(CH2)3
$$\begin{pmatrix}
R^{63} \\
Si \\
Si \\
R^{64} \\
M12
\end{pmatrix} CH3
$$CH3$$

$$CH3$$

$$CH3$$

$$CH3$$$$$$

—он

-continued
$$\begin{array}{c} \text{-continued} \\ \\ -\text{C} \\ \\ \text{C} \\ \\ \text{CH}_3 \\ \\ \text{CH}_3 \end{array}$$

In the formula (3-3), m¹² represents the number of repetition enclosed in brackets; the average of m12 in the polyester C ranges from 20 to 100; R⁶³ and R⁶⁴ each independently represent a methyl group or a phenyl group.

Hereinafter, specific examples of the terminal structure represented by the formula (3-3) are shown:

(3-3-1)

-continued

$$\begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \\ \\ \end{array} \\ \end{array} \\ \begin{array}{c} \begin{array}{c} \text{CH}_3 \\ \\ \text{Si} \end{array} \\ \begin{array}{c} \text{CH}_3 \\ \\ \text{Si} \end{array} \\ \begin{array}{c} \text{CH}_3 \\ \\ \text{I} \end{array} \\ \begin{array}{c} \text{CH}_3 \\ \\ \text{I} \end{array} \\ \begin{array}{c} \text{CH}_3 \\ \\ \text{I} \end{array} \\ \end{array} \end{array}$$

The polyesters having a siloxane bond can be used alone or in combination by mixing.

The content of the polyester having a siloxane bond in the emulsion can be not less than 0.01% by mass and not more than 5% by mass based on the total mass of the charge transporting substance and the binder resin. At a content within this range, the effect of stability of the emulsion by use of the polyester having a siloxane bond can be sufficiently obtained, and the effect of sufficient electrophotographic properties can be obtained.

Examples of the polystyrene having a siloxane bond include a polystyrene D having a repeating structural unit represented by the following formula (4-1) and a repeating structural unit represented by the following formula (4-2):

$$\begin{array}{c} \begin{array}{c} \begin{array}{c} \text{CH}_{3} \\ \end{array} \\ \begin{array}{c} \text{CH}_{2} \\ \end{array} \\ \begin{array}{c} \text{C} \\ \end{array} \\ \begin{array}{c} \text{C$$

where m⁷ represents an integer selected from 1 to 10; m⁸ ⁵⁰ represents an integer selected from 20 to 100.

Hereinafter, specific examples of the formula (4-1) are shown.

TABLE 4

Repeating structural unit represented by formula (4-1)	m ⁷	m ⁸	
Repeating structural unit example (4-1-1)	1	20	60
Repeating structural unit example (4-1-2)	3	20	
Repeating structural unit example (4-1-3)	3	40	
Repeating structural unit example (4-1-4)	1	60	65

TABLE 4-continued

Repeating structural unit represented by formula (4-1)	m^7	m ⁸
Repeating structural unit example (4-1-5)	3	100

The polystyrenes having a siloxane bond can be used 10 alone or in combination by mixing.

The content of the polystyrene having a siloxane bond in the emulsion can be not less than 0.5% by mass and not more than 10% by mass based on the total mass of the charge transporting substance and the binder resin. At a content within this range, the effect of stability of the emulsion by used of the polystyrene having a siloxane bond can be sufficiently obtained, and the effect of sufficient electrophotographic properties can be obtained.

Examples of the silicone oil include a compound represented by the following formula (5):

where R⁴² to R⁴⁵ each independently represent a methyl group or a phenyl group; m⁹ represents an integer selected from 20 to 100.

Hereinafter, specific examples of the silicone oil are shown:

(5-6)

(5-7) 10

-continued

$$\begin{array}{c} CH_{3} \\ H_{3}C - Si - O \\ CH_{3} \\ CH_{3} \end{array} \begin{array}{c} CH_{3} \\ Si - O \\ CH_{3} \\ CH_{3} \end{array} \begin{array}{c} CH_{3} \\ Si - O \\ CH_{3} \\ CH_{3} \end{array} \begin{array}{c} CH_{3} \\ CH_{3} \\ CH_{3} \end{array}$$

$$H_3C$$
 CH_3 $-$

The silicone oils can be used alone or in combination by mixing.

The content of the silicone oil in the emulsion can be not less than 0.5% by mass and not more than 10% by mass based on the total mass of the charge transporting substance and the binder resin. At a content within this range, the effect of stability of the emulsion by use of the silicone oil can be 30 sufficiently obtained, and the effect of sufficient electrophotographic properties can be obtained.

Examples of the polyolefin include aliphatic hydrocarbons

Hereinafter, specific examples of the polyolefin are $_{35}$ are shown: shown:

$$H_3C \leftarrow CH_2 \rightarrow {}_8CH_3$$
 (6-1)

$$H_3C - CH_2 - 10CH_3$$
 (6-2)

$$H_3C \leftarrow CH_2 \rightarrow_{14}CH_3$$
 (6-3)

$$H_3C \leftarrow CH_2 \rightarrow_{16}CH_3$$
 (6-4)

$$H_3C \leftarrow CH_2 \rightarrow_{22}CH_3$$
 (6-5)

$$H_3C+CH_2+_{30}CH_3$$
 (6-6)

$$H_3C \leftarrow CH_2 \rightarrow _{38}CH_3$$
 (6-7)

The polyolefins can be used alone or in combination by mixing.

The content of the polyolefin in the emulsion can be not less than 1% by mass and not more than 10% by mass based on the total mass of the charge transporting substance and the binder resin. At a content within this range, the effect of stability of the emulsion by use of the polyolefin can be 55 sufficiently obtained, and the effect of sufficient electrophotographic properties can be obtained.

Examples of the aliphatic acid, aliphatic acid amide, and aliphatic acid ester include a compound having a repeating structure represented by the following formula (7-1):

where R^{46} represents an alkyl group having 10 to 40 carbon atoms; R^{47} represents a hydrogen atom, an amino group and an alkyl group having 10 to 40 carbon atoms.

Hereinafter, specific examples of the aliphatic acid are shown:

$$C_{22}H_{45}$$
— \ddot{C} — O — H (7-1-6)

$$C_{30}H_{61}$$
— C — O — H

Hereinafter, specific examples of the aliphatic acid amide are shown:

$$C_{10}H_{21}$$
— C — O — NH_2 (7-1-9)

$$C_{14}H_{29}$$
 C O NH_2 (7-1-10)

$$C_{30}H_{61}$$
— \ddot{C} — O — NH_{2}
O
(7-1-13)

$$C_{40}H_{81}$$
 C_{-0} C_{-0} C_{-0}

Hereinafter, specific examples of the aliphatic acid ester are shown, but not limited to these:

$$\begin{array}{c} C_{10}H_{21} & C & (7-1-15) \\ \\ C_{10}H_{21} & C & (7-1-16) \\ \\ C_{14}H_{29} & C & (7-1-16) \\ \\ C_{18}H_{37} & C & (7-1-17) \\ \\ C_{18}H_{37} & C & (7-1-18) \\ \\ C_{25}H_{51} & C & (7-1-18) \\ \\ C_{25}H_{51} & C & (7-1-18) \\ \\ C_{30}H_{61} & C & (7-1-19) \\ \\ C_{30}H_{61} & C & (7-1-20) \\ \\ C_{40}H_{81} & C & (7-1-21) \\ \\ C_{40}H_{81} & C & (7-1-21) \\ \\ \end{array}$$

The aliphatic acids, aliphatic acid amides, and aliphatic acid esters can be used alone or in combination by mixing.

The content of the aliphatic acid, aliphatic acid amide, and aliphatic acid ester in the emulsion can be not less than 1% by mass and not more than 10% by mass based on the total mass of the charge transporting substance and the binder resin. At a content within this range, the effect of stability of the emulsion by use of the aliphatic acid, aliphatic acid amide, and aliphatic acid ester can be sufficiently obtained, and the effect of sufficient electrophotographic properties can be obtained.

The fluorine-atom-containing polyacrylate and fluorine-atom-containing polymethacrylate, the polycarbonate having a siloxane bond, the polystyrene having a siloxane bond, the silicone oil, the polyolefin, the aliphatic acid, aliphatic acid amide, and aliphatic acid ester can be used in combination by mixing.

A solvent used to prepare the solution containing the charge transporting substance and the compound that reduces the surface energy is those that dissolve the charge transporting substance. A liquid (hydrophobic solvent) whose solubility in water is 1.0% by mass or less at 25° C. and 1 atmosphere (atmospheric pressure) can be used.

Hereinafter, representative examples of the hydrophobic solvent are shown in table 5. The water solubility in table 5 means solubility in water at 25° C. and 1 atmospheric pressure (atmospheric pressure) which is indicated by % by 55 mass.

TABLE 5

Representative examples of hydrophobic solvent				
No	Name	Water solubility		
(E-1)	Toluene	0.1% by mass	_	
(E-2)	Chloroform	0.8% by mass		
(E-3)	o-Dichlorobenzene	0.0% by mass		
(E-4)	Chlorobenzene	0.1% by mass		
(E-5)	o-Xylene	0.0% by mass		

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TABLE 5-continued

Representative examples of hydrophobic solvent			
No	Name	Water solubility	
(E-6)	Ethylbenzene	0.0% by mass	
(E-7)	Phenetole	0.1% by mass	

Among these hydrophobic solvents, solvents having an aromatic ring structure are preferable, and at least one selected from the group consisting of toluene and xylene is more preferable from the viewpoint of stabilizing the emulsion. These hydrophobic solvents can be used in combination by mixing.

In the solution containing the charge transporting substance and the compound that reduces the surface energy, a hydrophilic solvent which is a solvent having solubility in water at 1 atmospheric pressure (atmospheric pressure) of 5.0% by mass or more can be mixed and used in addition of the hydrophobic solvent above.

Hereinafter, representative examples of the hydrophilic solvent are shown in Table 6. The water solubility in Table 6 means solubility in water at 25° C. and 1 atmospheric pressure (atmospheric pressure) which is indicated by % by mass

TABLE 6

Representative examples of hydrophilic solvent					
No	Name	Water solubility			
F-1	Tetrahydrofuran	100.0% by mass or more			
F-2	Dimethoxymethane	32.3% by mass			
F-3	1,2-Dioxane	100.0% by mass or more			
F-4	1,3-Dioxane	100.0% by mass or more			
F-5	1,4-Dioxane	100.0% by mass or more			
F-6	1,3,5-Trioxane	21.1% by mass			
F-7	Methanol	100.0% by mass or more			
F-8	2-Pentanone	5.9% by mass			
F-9	Ethanol	100.0% by mass or more			
F-10	Tetrahydropyran	100.0% by mass or more			
F-11	Diethylene glycol dimethyl ether	100.0% by mass or more			
F-12	Ethylene glycol dimethyl ether	100.0% by mass or more			
F-13	Propylene glycol n- butyl ether	6.0% by mass			
F-14	Propylene glycol monopropyl ether	100.0% by mass or more			
F-15	Ethylene glycol monomethyl ether	100.0% by mass or more			
F-16	Diethylene glycol monoethyl ether	100.0% by mass or more			
F-17	Ethylene glycol monoisopropyl ether	100.0% by mass or more			
F-18	Ethylene glycol monobutyl ether	100.0% by mass or more			
F-19	Ethylene glycol monoisobutyl ether	100.0% by mass or more			
F-20	Ethylene glycol monoallyl ether	100.0% by mass or more			
F-21	PROPYLENE GLYCOL MONOMETHYL ETHER	100.0% by mass or more			
F-22	Dipropylene glycol monomethyl ether	100.0% by mass or more			
F-23	Tripropylene glycol monomethyl ether	100.0% by mass or more			
F-24	Propylene glycol monobutyl ether	6.4% by mass			
F-25	Propylene glycol	20.5% by mass			
F-26	Diethylene glycol	100.0% by mass or more			
	methyl ethyl ether	,			

Representative examples of hydrophilic solvent					
No	Name	Water solubility			
F-27	Diethylene glycol diethyl ether	100.0% by mass or more			
F-28	Dipropylene glycol dimethyl ether	37.0% by mass			
F-29	Propylene glycol diacetate	7.4% by mass			
F-30	Methyl acetate	19.6% by mass			
F-31	Ethyl acetate	8.3% by mass			
F-32	n-Propyl alcohol	100.0% by mass or more			
F-33	3-Methoxy butanol	100.0% by mass or more			
F-34	3-Methoxybutyl acetate	6.5% by mass			
F-35	Ethylene glycol	100.0% by mass or more			

Among these hydrophilic solvents, ether solvents are preferable, and at least one selected from the group consisting of tetrahydrofuran and dimethoxymethane is more preferable from the viewpoint of stabilizing the emulsion.

monomethyl ether

acetate

These hydrophilic solvents can be used in combination by mixing. Particularly, in the case where a coat of the emulsion 25 is formed on the support by dip coating in the step of forming the coat of the emulsion on the support, use of a hydrophilic solvent having a relatively low boiling point of 100° C. or less is preferable. This is more preferable from the viewpoint of uniformity of the coat because the solvent 30 is quickly removed in the heating and drying step.

Next, a method of preparing the emulsion by dispersing the solution prepared by the method above in water will be described

As an emulsifying method for preparing an emulsion, 35 existing emulsifying methods can be used. The emulsion contains at least the charge transporting substance, the compound that reduces the surface energy, and the binder resin in the emulsion particles in the state where the charge transporting substance, the compound that reduces the surface energy, and the binder resin are partially or entirely dissolved in the emulsion particles. Hereinafter, as specific emulsifying methods, a stirring method and a high pressure collision method will be shown, but the production method according to the present invention will not be limited to 45 these.

The stirring method will be described. In this method, the charge transporting substance, the compound that reduces the surface energy, and the binder resin are dissolved in the solvent (hydrophobic solvent, hydrophilic solvent) to prepare a solution. The solution is mixed with water, and stirred by a stirrer. Here, from the viewpoint of the electrophotographic properties, water can be ion exchange water from which metal ions and the like are removed with an ion exchange resin or the like. The ion exchange water can have 55 a conductivity of 5 μ S/cm or less. As the stirrer, a stirrer enabling high speed stirring can be used because a uniform emulsion can be prepared in a short time. Examples of the stirrer include a homogenizer (Physcotron) made by MICROTEC CO., LTD. and a circulation homogenizer 60 (Cleamix) made by M Technique Co., Ltd.

The high pressure collision method will be described. In this method, the charge transporting substance, the compound that reduces the surface energy, and the binder resin are dissolved in the solvent (hydrophobic solvent, hydrophilic solvent) to prepare a solution. The solution is mixed with water, and the mixed solution is collided under high 28

pressure. Thus, an emulsion can be prepared. Alternatively, without mixing the solution with water, the solution may be collided with water as individual solutions to prepare an emulsion. Examples of a high pressure colliding apparatus include a Microfluidizer M-110EH made by Microfluidics Corporation in U.S. and a Nanomizer YSNM-2000AR made by YOSHIDA KIKAI CO., LTD.

As the mixing ratio of water to the solution containing the charge transporting substance, the compound that reduces the surface energy, and the binder resin in the emulsion, water/solution is 3/7 to 8/2, and can be 5/5 to 7/3 from the viewpoint of obtaining an emulsion having a high concentration of the solid content while stability of the emulsion is kept.

The ratio of water to the solvent (hydrophobic solvent, hydrophilic solvent) can be 4/6 to 8/2 (water has a higher proportion) from the viewpoint of reducing the size of the oil droplet in emulsifying and stabilizing the emulsion. The ratio above can be adjusted in the range in which the charge transporting substance and the binder resin are dissolved in an organic solvent. Thus, the size of the oil droplet is reduced to enhance solution stability.

In the oil droplets in the emulsion, the proportion of the charge transporting substance, the compound that reduces the surface energy, and the binder resin to the solvent can be 10 to 50% by mass. The proportion of the charge transporting substance to the binder resin to be contained in the solution is preferably in the range of 4:10 to 20:10 (mass ratio), and more preferably in the range of 5:10 to 12:10 (mass ratio).

Moreover, the emulsion may contain a surfactant for the purpose of further stabilizing the emulsion. As the surfactant, a nonionic surfactant (nonionic surfactant) can be used from the viewpoint of suppressing reduction in the electrophotographic properties. The nonionic surfactant has a hydrophilic portion which is a non-electrolyte, that is, not ionized. Examples of the nonionic surfactant include:

NAROACTY Series, EMULMIN Series, SANNONIC Series, and NEWPOL Series made by Sanyo Chemical Industries, Ltd., EMULGEN Series, RHEODOL Series, and EMANON Series made by Kao Corporation,

Adekatol Series, ADEKA ESTOL Series, and ADEKA NOL Series made by ADEKA Corporation, and nonionic surfactant Series among Newcol Series made by NIPPON NYUKAZAI CO., LTD.

Surfactants above can be used alone or in combination. The surfactant having an HLB value (Hydrophile-Lipophile Balance value) in the range of 8 to 15 can be selected for stabilization of the emulsion.

The amount of the surfactant to be added is preferably as small as possible from the viewpoint of preventing reduction in the electrophotographic properties. The content of the surfactant in the emulsion is preferably in the range of 0% by mass to 1.5% by mass, and more preferably in the range of 0% by mass to 0.5% by mass based on the total mass of the charge transporting substance and the binder resin. The surfactant may be contained in water, or may be contained in the solution containing the charge transporting substance the compound that reduces the surface energy, and the binder resin. Alternatively, the surfactant may be contained in both water and the solution.

Moreover, the emulsion may contain an antifoaming agent, a viscoelastic adjuster and the like in the range in which the effect of the present invention is not inhibited.

The average particle diameter of the emulsion particle in the emulsion is preferably in the range of 0.1 to $20.0 \, \mu m$, and

more preferably in the range of 0.1 to 5.0 μm from the viewpoint of stability of the emulsion.

Next, a method of applying the coat of the emulsion onto a support will be described.

As a step of forming the coat of the emulsion on the support, any of existing coating methods such as a dip coating method, a ring coating method, a spray coating method, a spinner coating method, a roller coating method, a Meyer bar coating method, and a blade coating method can be used. From the viewpoint of productivity, the dip coating can be used. According to the dip coating method, the emulsion can be applied onto a support to form a coat.

Next, a step of heating the coat to form a charge transporting layer will be described. The formed coat is heated to form a charge transporting layer.

The coat of the emulsion may be formed on the charge generating layer. Alternatively, the coat of the emulsion may be formed on an undercoat layer, and the charge generating layer may be formed on the coat. Further, in the case where 20 the charge transporting layer has a laminate structure (first charge transporting layer, second charge transporting layer), the coat of the emulsion may be formed on the first charge transporting layer to form the second charge transporting layer. Alternatively, using the coat of the emulsion, both of 25 the first charge transporting layer and the second charge transporting layer may be formed.

In the present invention, the emulsion containing at least the charge transporting substance, the compound that reduces the surface energy, and the binder resin is applied to 30 form the coat. For this reason, by heating the coat, the dispersion medium (water) can be removed and the emulsion particles can be brought into close contact with each other at the same time. Thereby, a more uniform coat can be formed. Thereby, a coat having high uniformity can be 35 formed. Further, if the emulsion particle has a smaller particle diameter, a film thickness having high uniformity can be quickly obtained after the dispersion medium is removed. Accordingly, a smaller particle diameter of the emulsion particle is preferable. A heating temperature can be 40 100° C. or more. Further, from the viewpoint of enhancing close contact of the emulsion particles, the heating temperature can be a heating temperature of the melting point or more of the charge transporting substance having the lowest melting point among the charge transporting substances that 45 form the charge transporting layer. By heating at a temperature of the melting point or more of the charge transporting substance, the charge transporting substance is fused. The binder resin is dissolved in the fused charge transporting substance. Thereby, a highly uniform coat can be formed. 50 layer. Further, heating can be performed at a heating temperature 5° C. or more higher than the melting point of the charge transporting substance having the lowest melting point among the charge transporting substances that form the charge transporting layer. Moreover, the heating temperature 55 can be 200° C. or less. Occurrence of modification or the like of the charge transporting substance can be suppressed, obtaining sufficient electrophotographic properties.

The film thickness of the charge transporting layer produced by the production method according to the present $_{60}$ invention is preferably not less than 3 μm and not more than 50 μm , and more preferably not less than 5 μm and not more than 35 μm .

Next, the configuration of the electrophotographic photosensitive member produced by the production method of 65 the electrophotographic photosensitive member according to the present invention above will be described.

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A cylindrical electrophotographic photosensitive member formed of a cylindrical support and a photosensitive layer (charge generating layer, charge transporting layer) formed thereon is usually widely used, but the electrophotographic photosensitive member can have a belt-like shape or a sheet-like shape, for example.

As the support, those having conductivity (electrically conductive support) can be used. A metallic conductive support made of aluminum, an aluminum alloy, stainless steel, or the like can be used. In the case of the aluminum or aluminum alloy conductive support, an ED tube, an EI tube, or those subjected to machining, electrochemical mechanical polishing, a wet or dry honing treatment can also be used. Moreover, a metallic conductive support or a resin conductive support having a layer of a coat formed by vacuum depositing aluminum, an aluminum alloy or an indium oxide-tin oxide alloy can also be used. Moreover, a conductive support formed by impregnating conductive particles such as carbon black, tin oxide particles, ittanium oxide particles, and silver particles into a resin, or a plastic having a conductive resin can also be used.

The surface of the support may be subjected to a machining treatment, a surface roughening treatment, an anodic oxidation treatment, or the like.

An electrically conductive layer may be provided between the support and an undercoat layer or charge generating layer described later. The electrically conductive layer can be obtained by forming a coat on the support using a coating solution for an electrically conductive layer in which conductive particles are dispersed in a resin, and drying the coat. Examples of the conductive particles include carbon black, acetylene black, metal powders of aluminum, nickel, iron, nichrome, copper, zinc, and silver, and metal oxide powders of conductive tin oxide and ITO.

Examples of the resin include polyester resins, polycarbonate resins, polyvinyl butyral, acrylic resins, silicone resins, epoxy resins, melamine resins, urethane resins, phenol resins and alkyd resins.

Examples of a solvent used in the coating solution for an electrically conductive layer include ether solvents, alcohol solvents, ketone solvents and aromatic hydrocarbon solvents.

The film thickness of the electrically conductive layer is preferably not less than 0.2 μm and not more than 40 μm , more preferably not less than 1 μm and not more than 35 μm , and still more preferably not less than 5 μm and not more than 30 μm .

An undercoat layer may be provided between the support or electrically conductive layer and the charge generating layer.

The undercoat layer can be formed by forming a coat on the support or electrically conductive layer using a coating solution for an undercoat layer having a resin, and drying or curing the coat.

Examples of the resin for the undercoat layer include polyacrylic acids, methyl cellulose, ethyl cellulose, polyamide resins, polyamide resins, polyamide resins, polyamide resins, polyumic acid resins, melamine resins, epoxy resins, polyurethane resins, and polyolefin resins. As the resin used for the undercoat layer, thermoplastic resins can be used. Specifically, thermoplastic polyamide resins or polyolefin resins can be used. As the polyamide resins, copolymerized nylons having low crystallinity or non-crystallinity and allowing application in a liquid state can be used. As the polyolefin resins, those in a state where those can be used as a particle dispersion liquid can be used. Further, polyolefin resins can be dispersed in an aqueous medium.

The film thickness of the undercoat layer is preferably not less than 0.05 μm and not more than 30 μm , and more preferably not less than 1 μm and not more than 25 μm . Moreover, the undercoat layer may contain a metal-oxide particle.

Moreover, the undercoat layer may contain a semi-conductive particle, an electron transporting substance, or an electron receiving substance.

A charge generating layer can be provided on the support, the electrically conductive layer or the undercoat layer.

Examples of the charge generating substance used in the electrophotographic photosensitive member include azo pigments, phthalocyanine pigments, indigo pigments and perylene pigments. These charge generating substances may be used alone or in combination. Among these, particularly metal phthalocyanines such as oxytitanium phthalocyanine, hydroxy gallium phthalocyanine, and chlorogallium phthalocyanine have high sensitivity and can be used.

Examples of a binder resin used in the charge generating 20 layer include polycarbonate resins, polyester resins, butyral resins, polyvinylacetal resins, acrylic resins, vinyl acetate resins and urea resins. Among these, particularly butyral resins can be used. These can be used alone, or can be used in combination by mixing or as a copolymer.

The charge generating layer can be formed by forming a coat using a coating solution for a charge generating layer obtained by dispersing the charge generating substance together with a binder resin and a solvent, and heating the coat. Alternatively, the charge generating layer may be a 30 deposited film of the charge generating substance.

Examples of a dispersing method include methods using a homogenizer, ultrasonic waves, a ball mill, a sand mill, an Attritor, and a roll mill.

The proportion of the charge generating substance to the 35 binder resin is preferably in the range of 1:10 to 10:1 (mass ratio), and particularly more preferably in the range of 1:1 to 3:1 (mass ratio).

Examples of the solvent used in the coating solution for a charge generating layer include alcohol solvents, sulfoxide 40 solvents, ketone solvents, ether solvents, ester solvents or aromatic hydrocarbon solvents.

The film thickness of the charge generating layer is preferably not less than 0.01 μm and not more than 5 μm , and more preferably not less than 0.1 μm and not more than 2 $_{45}$ μm .

Moreover, a variety of a sensitizer, an antioxidant, an ultraviolet absorbing agent, a plasticizer and the like can also be added to the charge generating layer when necessary. In order to prevent stagnation of a flow of charges in the charge generating layer, an electron transporting substance or electron receiving substance may be contained in the charge generating layer.

The charge transporting layer is provided on the charge generating layer.

The charge transporting layer is produced by the production method above.

Deterioration preventing materials such as an antioxidant, an ultraviolet absorbing agent, and a light stabilizer, and fine particles such as organic fine particles and inorganic fine 60 particles may be added to each of the layers in the electrophotographic photosensitive member. Examples of the antioxidant include hindered phenol antioxidants, hindered amine light stabilizers, sulfur atom-containing antioxidants, and phosphorus atom-containing antioxidants. Examples of 65 the organic fine particles include molecule resin particles such as fluorine atom-containing resin particles, polystyrene

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fine particles, and polyethylene resin particles. Examples of the inorganic fine particles include metal oxides such as silica and alumina.

In application of the coating solutions for the respective layers above, coating methods such as a dip coating method, a spray coating method, a spinner coating method, a roller coating method, a Meyer bar coating method, and a blade coating method can be used.

Moreover, a shape of depressions and projections (a shape of depressions, a shape of projections) may be formed on the surface of the charge transporting layer which is a surface layer in the electrophotographic photosensitive member. As a method of forming a shape of depressions and projections, a known method can be used. Examples of the forming method include a method for forming a shape of depressions by spraying polished particles to the surface, a method for forming a shape of depressions and projections by bringing a mold having a shape of depressions and projections into contact with the surface under pressure, and a method for forming a shape of depressions by irradiating the surface with laser light. Among these, a method can be used in which a mold having a shape of depressions and projections is brought into contact with the surface of the surface layer of the electrophotographic photosensitive member under pressure to form a shape of depressions and projections.

FIG. 2 shows an example of a schematic configuration of an electrophotographic apparatus including a process cartridge having the electrophotographic photosensitive member according to the present invention.

In FIG. 2, a cylindrical electrophotographic photosensitive member 1 is shown. The electrophotographic photosensitive member 1 is rotated and driven around a shaft 2 in the arrow direction at a predetermined circumferential speed. The surface of the electrophotographic photosensitive member 1 rotated and driven is uniformly charged at a positive or negative potential by a charging unit (primary charging unit: charging roller or the like) 3. Next, the surface of the electrophotographic photosensitive member 1 receives expositing light (image expositing light) 4 output from an exposing unit (not shown) such as slit exposure and laser beam scanning exposure. Thus, an electrostatic latent image corresponding to a target image is sequentially formed on the surface of the electrophotographic photosensitive member 1

The electrostatic latent image formed on the surface of the electrophotographic photosensitive member 1 is developed with a toner included in a developer in a developing unit 5 to form a toner image. Next, the toner image carried on the surface of the electrophotographic photosensitive member 1 is sequentially transferred onto a transfer material (such as paper) P by a transfer bias from a transferring unit (transfer roller or the like) 6. The transfer material P is extracted from a transfer material feeding unit (not shown) and fed to a region between the electrophotographic photosensitive member 1 and the transferring unit 6 (contact region) in synchronization with the rotation of the electrophotographic photosensitive member 1.

The transfer material P to which the toner image is transferred is separated from the surface of the electrophotographic photosensitive member 1, and introduced to a fixing unit 8 to fix the image. Thereby, the transfer material P is printed out to the outside the apparatus as an image forming product (print, copy).

The surface of the electrophotographic photosensitive member 1 after transfer of the toner image is cleaned by removing a transfer remaining developer (toner) by a cleaning unit (cleaning blade or the like) 7. Next, the surface of

the electrophotographic photosensitive member 1 is discharged by a pre-expositing light (not shown) from a pre-exposing unit (not shown), and repeatedly used for formation of an image. As shown in FIG. 2, in the case where the charging unit 3 is a contact charging unit using a charging of 15 roller, pre-exposure is not always necessary.

Among the components such as the electrophotographic photosensitive member 1, the charging unit 3, the developing unit 5, the transferring unit 6 and the cleaning unit 7, a plurality of the components may be accommodated in a container and integrally formed into a process cartridge, and the process cartridge may be formed attachably to and detachably from the main body of the electrophotographic apparatus such as a copier and a laser beam printer. In FIG. 2, the electrophotographic photosensitive member 1, the charging unit 3, the developing unit 5 and the cleaning unit 7 are integrally supported and formed as a cartridge, and the cartridge is formed as a process cartridge 9 attachably to and detachably from the main body of the electrophotographic apparatus using a guiding unit 10 such as a rail in the main body of the electrophotographic apparatus.

EXAMPLES

Hereinafter, the present invention will be described more 25 in detail using Examples and Comparative Examples. The present invention will not be limited by Examples below. In Examples, "parts" mean "parts by mass."

Example 1

(Preparation of Emulsion)

5 parts of the compound represented by the formula (CTM-1) and 5 parts of the compound represented by the formula (CTM-7) as the charge transporting substance, and 35 10 parts of a polycarbonate resin having a repeating structural unit represented by the formula (B1-1) (weight average molecular weight Mw=57,000), and 0.1 parts of the compound represented by the formula (1-2) as the binder resin were dissolved in 60 parts of toluene to prepare a solution. 40 Next, while 120 parts of ion exchange water (conductivity of 0.2 μS/cm) was stirred by a homogenizer (Physcotron) made by MICROTEC CO., LTD. at a rate of 3,000 turns/min, 80.1 parts of the solution was gradually added for 10 minutes. After dropping was completed, the number of rotation of the 45 homogenizer was raised to 7,000 turns/min and stirring was performed for 20 minutes. Then, the obtained solution was emulsified by a high pressure collision dispersing machine Nanomizer (made by YOSHIDA KIKAI CO., LTD.) on a pressure condition of 150 MPa to obtain an emulsion (80.1 50 parts).

(Evaluation of Solution Stability of Emulsion)

After the emulsion was prepared according to the method above, the emulsion was visually evaluated and the particle diameter of the emulsion particle was evaluated. Further, the 55 prepared emulsion was left as it was for 2 weeks (under an environment of the temperature of 25° C. and the humidity of 50% RH). After the state of the emulsion after leaving was observed, the emulsion was stirred at a rate of 1,000 turns/min for 3 minutes using a homogenizer made by MICRO-TEC CO., LTD. The state of the emulsion after stirring was visually observed in the same manner. The average particle diameters of the emulsion particle in the emulsion before and after leaving the emulsion as it was and stirring it were measured. In the measurement of the average particle diameter of the emulsion particle, the emulsion was diluted with water, and the average particle diameter was measured using

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an ultracentrifugal automatic particle size distribution analyzer (CAPA700) made by HORIBA, Ltd. The results are shown in Table 14. The states of the emulsion obtained in Example 1 before and after leaving were not greatly changed even by visually observation. The average particle diameter hardly changed, and the emulsion was kept stably. The results of evaluation are shown in Table 7.

Examples 2 to 296

Emulsions were prepared by the same method as that in Example 1 except that the kinds and ratios of the charge transporting substance, the compound that reduced the surface energy, the binder resin, and the solvent were changed as shown in Table 7 to Table 13. The results of evaluation of solution stability of the obtained emulsions are shown in Tables 14 to 15. In Examples 5, 15, 45, 58, 105, 118, 144, 155, 173, 185, 202, 215, 236, and 242, 0.5% by mass of a surfactant (trade name: NAROACTY CL-85, made by Sanyo Chemical Industries, Ltd., HLB=12.6) was further contained based on the total mass of the charge transporting substance and the binder resin.

Example 297

An emulsion was prepared by the same method as that in Example 3 except that in Example 3, the fluorine-containing acrylate used in Example 6 and the silicone oil used in Example 173 were mixed and used. The results of evaluation of solution stability of the obtained emulsion are shown in Table 15.

Examples 298 to 300

Emulsions were prepared by the same method as that in Example 297 except that in Example 297, the fluorine-containing acrylate used in Example 6 was replaced by the compound shown below. The results of evaluation of solution stability of the obtained emulsions are shown in Table 15. In Example 298, the fluorine-containing acrylate used in Example 6 was replaced by the polycarbonate A used in Example 36. In Example 299, the fluorine-containing acrylate used in Example 6 was replaced by the polyester C used in Example 98. In Example 300, the fluorine-containing acrylate used in Example 6 was replaced by the polystyrene D used in Example 139.

Example 701

An emulsion was prepared by the same method as that in Example 36 except that in Example 36, the hydrophobic solvent was replaced by (E-7). The solution stability of the obtained emulsion is shown in Table 15.

Comparative Example 1

An emulsion containing a charge transporting substance and a binder resin was prepared according to the method described in Japanese Patent Application Laid-Open No. 2011-128213 as follows.

5 parts of the compound represented by the formula (CTM-7) as the charge transporting substance, and 5 parts of a polycarbonate resin having a repeating structural unit represented by the formula (B1-1) (weight average molecular weight Mw=36,000) as the binder resin were dissolved in 40 parts of toluene to prepare the solution (50 parts). Next, 1.5 parts of a surfactant (trade name: NAROACTY CL-70

made by Sanyo Chemical Industries, Ltd.) was added to 48.5 parts of water. While the water was stirred at a rate of 3,000 turns/min with a homogenizer made by MICROTEC CO., LTD., the solution was added, and stirred for 10 minutes. Further, the number of rotation of the homogenizer made by MICROTEC CO., LTD. was raised to 7,000 turns/min and stirring was performed for 20 minutes. Then, the obtained solution was emulsified on a pressure condition of 150 MPa using a high pressure collision dispersing machine Nanomizer (made by YOSHIDA KIKAI CO., LTD.) to obtain 100 parts of an emulsion. In the obtained emulsion, the states of the emulsion and the average particle diameters before leaving and after leaving and stirring with a homogenizer, were measured by the same method as that in Example 1. The results are shown in Table 16.

In the state after leaving of the emulsion obtained in Comparative Example 1, sediment of the oil droplet component was found, and the oil droplet component partially coalesced and aggregates were found on the bottom. Unlike the emulsion immediately after the emulsion was prepared, 20 in the emulsion after stirring, aggregation of the oil droplet component was found, and the state of an emulsion having high uniformity could not be obtained.

Comparative Example 2

An emulsion was prepared by the same method as that in Comparative Example 1 except that in Comparative Example 1, a compound represented by the formula (CTM-3) was used as the charge transporting substance, and 30 chlorobenzene was used as the solvent. The stability of the obtained emulsion for a charge transporting layer was evaluated by the same method as that in Comparative Example 1. The results are shown in Table 16.

Comparative Example 3

An emulsion was prepared by the same method as that in Comparative Example 1 except that in Comparative

Example 1, 20 parts of chlorobenzene was replaced by 20 parts of chloroform as the solvent, and the surfactant was replaced by NAROACTY CL-85 made by Sanyo Chemical Industries, Ltd. The stability of the obtained emulsion was evaluated by the same method as that in Comparative Example 1. The results are shown in Table 16.

Comparative Example 4

An emulsion was prepared by the same method as that in Comparative Example 1 except that in Comparative Example 1, 20 parts of chlorobenzene was replaced by 20 parts of o-dichlorobenzene as the solvent, and the surfactant was replaced by EMULMIN 140 made by Sanyo Chemical Industries, Ltd. The stability of the obtained emulsion was evaluated by the same method as that in Comparative Example 1. The results are shown in Table 16.

Comparative Example 5

An emulsion was prepared by the same method as that in Comparative Example 1 except that in Comparative Example 1, zinc stearate was further contained. The stability of the obtained emulsion was evaluated by the same method as that in Comparative Example 1. The results are shown in Table 16.

Comparative Example 6

An emulsion was prepared by the same method as that in Comparative Example 1 except that in Comparative Example 1, zinc linolenate was further contained. The stability of the obtained emulsion was evaluated by the same method as that in Comparative Example 1. The results are shown in Table 16.

TABLE 7

	Fluorine-atom- acrylate, fluori			Binder resin	and ratio	-	
	containing met	hacrylate	_		Weight	Kind and ratio	of solvent
Example	Repeating structural unit, ratio	Content (%)	Charge transporting substance and ratio	Repeating structural unit, ratio	average molecular weight	Hydrophobic solvent/hydrophilic solvent, ratio	Water/solvent
1	(1-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)	6/4
2	(1-3)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)/(F-2) = 9/1	6/4
3	(1-10)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/(F-1) = 9/1	6/4
4	(1-11)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-5)	6/4
5	(1-2)/(1-10) = 7/3	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/(F-1) = 9/1	6/4
6	(1-2)/(1-10) = 5/5	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/(F-1) = 9/1	6/4
7	(1-2)/(1-10) = 3/7	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/(F-1) = 9/1	6/4
8	(1-2)/(1-10) = 5/5	0.5%	CTM-1	(B1-1)	57000	(E-5)	6/4
9	(1-2)/(1-10) = 5/5	0.5%	CTM-1/CTM-7 = 7/3	(B1-1)	57000	(E-5)/(F-2) = 9/1	6/4
10	(1-2)/(1-10) = 5/5	0.5%	CTM-1/CTM-7 = 3/7	(B1-1)	57000	(E-4)/(F-1) = 9/1	6/4
11	(1-2)/(1-10) = 5/5	0.5%	CTM-7	(B1-1)	57000	(E-4)/(F-2) = 9/1	6/4
12	(1-2)/(1-10) = 5/5	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	14000	(E-1)/(F-2) = 9/1	6/4
13	(1-2)/(1-10) = 5/5	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	120000	(E-1)/(F-1) = 9/1	6/4

TABLE 7-continued

	Fluorine-atom-co acrylate, fluoring			Binder resin	and ratio	_	
	containing meth	acrylate	_		Weight	Kind and ratio	of solvent
Example	Repeating structural unit, ratio	Content (%)	Charge transporting substance and ratio	Repeating structural unit, ratio		Hydrophobic solvent/hydrophilic solvent, ratio	Water/solvent
14	(1-2)/(1- 10) = 5/5	0.5%	CTM-1/CTM-7 = 5/5	(B1-2)	55000	(E-5)	6/4
15	(1-2)/(1-10) = 5/5	0.5%	CTM-1/CTM-7 = 5/5	(B1-3)	53000	(E-4)/(F-1) = 9/1	6/4
16	(1-2)/(1-10) = 5/5	0.5%	CTM-1/CTM-7 = 5/5	(B1-2)/(B1-3) = 3/7	55000	(E-5)/(F-1) = 9/1	6/4
17	(1-2)/(1-10) = 5/5	0.5%	CTM-1/CTM-7 = 5/5	(B1-2)/(B1-3) = 5/5	55000	(E-1)/(F-1) = 9/1	6/4
18	(1-2)/(1-10) = 5/5	0.5%	CTM-1/CTM-7 = 5/5	(B1-2)/(B1-3) = 7/3	55000	(E-4)/(F-2) = 9/1	6/4
19	(1-2)/(1-10) = 5/5	0.5%	CTM-1/CTM-7 = 5/5	(B2-1)	120000	(E-1)	6/4
20	(1-2)/(1-10) = 5/5	0.5%	CTM-1/CTM-7 = 5/5	(B2-2)	120000	(E-5)/(F-2) = 9/1	6/4
21	(1-2)/(1-10) = 5/5	0.5%	CTM-1/CTM-7 = 5/5	(B2-1)/(B2-2) = 7/3	120000	(E-1)/(F-2) = 9/1	6/4
22	(1-2)/(1-10) = 5/5	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)	6/4
23	(1-2)/(1-10) = 5/5	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/(F-1) = 9/1	6/4
24	(1-2)/(1-10) = 5/5		CTM-1/CTM-7 = 5/5	(B1-1)	57000	(F-1)	6/4
25	(1-1)/(1-2) = 5/5		CTM-2	(B1-4)	57000	(E-2)/(F-1) = 9/1	6/4
26	(1-4)/(1-5) = 5/5	1%	CTM-3	(B1-6)	57000	(E-6)/(F-10) = 9/1	6/4
27	(1-5)/(1-11) = 5/5	5%	CTM-4	(B1-8)	57000	(E-3)/(F-21) = 9/1	6/4
28	(1-6)/(1-3) = 5/5	0.3%	CTM-5	(B1-5)/ (B1-7) = 5/5	57000	(E-2)/(F-32) = 9/1	6/4
29	(1-7)/(1-1) = 5/5	0.5%	CTM-6	(B2-3)	57000	(E-2)/(F-20) = 9/1	6/4
30	(1-8)/(1-4) = 5/5		CTM-8	(B2-5)	57000	(E-6)/(F-11) = 9/1	6/4
31	(1-9)/(1-7) = 5/5		CTM-9	(B2-6)	57000	(E-6)/(F-7) = 9/1	6/4
32	(1-12)/ (1-10) = 5/5	0.3%	CTM-1/CTM-5 = 7/3	(B2-4)/ (B2-6) = 5/5	57000	(E-6)/(F-16) = 9/1	6/4
33	(1-3)/(1-6) = 5/5	0.1%	CTM-1/CTM-5 = 5/5	(B2-2)/ (B2-4) = 5/5	57000	(E-6)/(F-5) = 9/1	6/4
34	(1-12)/ (1-14) = 5/5	1%	CTM-1/CTM-5 = 3/7	(B2-3)/ (B2-5) = 5/5	57000	(E-3)/(F-3) = 9/1	6/4

TABLE 8

							Kind and ratio o	of solvent
	Polycarbonate ha	wing siloxane bond		Charge transporting	Binder	resin	Hydrophobic solvent/hydro-	
Example	Repeating structural unit, ratio	Terminal structure	Content (%)	substance and ratio	Repeating structural unit, ratio	Weight average molecular weight		Water/ solvent
35	(2-1-1)/(B1-1) = 9/1	_	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)	6/4
36	(2-1-2)/(B1-1) = 9/1	_	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/ (F-1) = 9/1	6/4
37	(2-1-2)/(B1-2) = 9/1	_	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-5)/ (F-1) = 9/1	6/4
38	(2-1-2)/(B1-2) = 9/1	_	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)	6/4
39	(2-1-2)/(2-1-6)/(B1-1) = 3.5/3.5/3	_	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/ (F-1) = 9/1	6/4
40	(2-1-2)/(2-1-6)/(B1-1) = 2.5/2.5/5	_	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/ (F-1) = 9/1	6/4
41	(2-1-2)/(2-1-6)/(B1-1) = 1.5/1.5/7	_	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/(F-1) = 9/1	6/4
42	(2-1-3)/(B1-1) = 9/1	_	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-5)/ (F-2) = 9/1	6/4
43	(2-1-4)/(B1-1) = 9/1	_	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-5)	6/4
44	(2-1-5)/(B1-1) = 9/1	_	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/ (F-1) = 9/1	6/4
45	(2-1-6)/(B1-1) = 9/1	_	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)	6/4

TABLE 8-continued

							Kind and ratio	of solven
	Polycarbonate 1	naving siloxane bon	d	Charge _transporting	Binder	resin	Hydrophobic _solvent/hydro-	
Example	Repeating structural unit, ratio	Terminal structure	Content (%)	substance and ratio	Repeating structural unit, ratio	Weight average molecular weight		Water/ solven
46	(2-1-7)/(B1-1) = 9/1	_	0.5%	CTM-1/CTM-7 =	(B1-1)	57000	(E-4)/	6/4
47	(2-2-1)/(B1-1) = 7/3	(2-4-2)	0.5%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-1) = 9/1 (E-5)	6/4
48	(2-2-2)/(B1-1) = 5/5	(2-4-2)	0.5%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(E-5)/	6/4
49	(2-2-3)/(B1-1) = 8/2	(2-4-2) · (2-5)	0.5%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-1) = 9/1 (E-4)/	6/4
50	(2-2-4)/(B1-1) = 7/3	(2-4-2)	0.5%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-1) = 9/1 (E-5)	6/4
51	(2-2-5)/(B1-1) = 7/3	_	0.5%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(E-5)/	6/4
52	(2-2-6)/(B1-1) = 7/3	(2-4-2)	0.5%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-1) = 9/1 (E-1)/	6/4
53	(2-2-7)/(B1-1) = 8/2	_	0.5%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-2) = 9/1 (E-4)/	6/4
54	(2-2-8)/(B1-1) = 8/2	(2-4-2)	0.5%		(B1-1)	57000	(F-2) = 9/1 (E-5)	6/4
55	(2-2-9)/(B1-1) = 5/5	_	0.5%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(E-1)/	6/4
56	(2-2-10)/(B1-1) = 6/4	_	0.5%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-2) = 9/1 (E-4)	6/4
57	(2-2-11)/(B1-1) = 8/2	(2-4-2)	0.5%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(E-1)	6/4
58	(2-2-12)/(B1-1) = 7/3	(2-4-2) · (2-6)	0.5%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(E-5)/	6/4
59	(2-2-13)/(B1-1) = 5/5	_	0.5%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-1) = 9/1 (E-5)	6/4
60	(2-2-14)/(B1-1) = 8/2	(2-4-2)	0.5%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(E-1)	6/4
61	(2-2-15)/(B1-1) = 7/3	_	0.5%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(E-5)	6/4
62	(2-2-16)/(B1-1) = 6/4	(2-4-2)	0.5%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(E-5)	6/4
63	(2-1-2)/(B1-1) = 9/1	_	0.5%	5/5 CTM-1	(B1-1)	57000	(E-4)/	6/4
64	(2-1-2)/(B1-1) = 9/1	_	0.5%	CTM-1/CTM-7 =	(B1-1)	57000	(F-1) = 9/1 (E-1)/	6/4
65	(2-1-2)/(B1-1) = 8/2	_	0.5%	7/3	,	57000	(F-2) = 9/1 (E-4)/	6/4
66	(2-1-2)(B1-1) = 9/1	_	0.5%	3/7 CTM-7	(B1-1)	57000	(F-1) = 9/1 (E-1)	6/4
67	(2-1-2)/(B1-1) = 9/1	_	0.5%	CTM-1/CTM-7 = 5/5		14000	(E-4)/ (F-2) = 9/1	6/4
68	(2-1-2)/(B1-1) = 9/1	_	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	120000	(E-4)	6/4
69	(2-1-2)/(B1-1) = 9/1	_	0.5%	CTM-1/CTM-7 = 5/5	(B1-2)	55000	(E-1)	6/4
70	(2-1-2)/(B1-1) = 9/1	_	0.5%	CTM-1/CTM-7 =	(B1-3)	53000	(E-5)/	6/4
71	(2-1-2)/(B1-1) = 9/1	_	0.5%	5/5 CTM-1/CTM-7 =	' ' '	55000	(F-2) = 9/1 (E-5)/	6/4
72	(2-1-2)/(B1-1) = 8/2	_	0.5%	5/5 CTM-1/CTM-7 =		55000	(F-2) = 9/1 (E-4)	6/4
73	(2-1-2)/(B1-1) = 9/1	_	0.5%	5/5 CTM-1/CTM-7 =		55000	(E-4)/	6/4
74	(2-1-2)/(B1-1) = 9/1	_	0.5%	5/5 CTM-1/CTM-7 =	7/3 (B2-1)	120000	(F-2) = 9/1 (E-5)/	6/4
75	(2-1-2)/(B1-1) = 9/1	_	0.5%	5/5 CTM-1/CTM-7 =	(B2-2)	120000	(F-2) = 9/1 (E-4)	6/4
76	(2-1-2)/(B1-1) = 9/1	_		5/5 CTM-1/CTM-7 =	,	120000	(E-1)/	6/4
77	(2-1-2)/(B1-1) = 9/1 (2-1-2)/(B1-1) = 9/1			5/5 CTM-1/CTM-7 =	7/3	57000	(F-2) = 9/1 (E-1)	6/4
	, , , ,	_		5/5	,			
78	(2-1-2)/(B1-1) = 9/1	_		CTM-1/CTM-7 = 5/5	,	57000	(E-4)/ (F-2) = 9/1	6/4
79	(2-1-2)/(B1-1) = 9/1	_	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(F-1)	6/4
80	(2-1-2)/(B1-1) = 9/1	_	2%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/ (F-1) = 9/1	6/4
81	(2-1-2)/(B1-1) = 9/1	_	2%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)/ (F-2) = 9/1	6/4

TABLE 8-continued

							Kind and ratio	of solvent
	Polycarbonate ha	aving siloxane bond		Charge _transporting	Binder	resin	Hydrophobic _solvent/hydro-	
Example	Repeating structural unit, ratio	Terminal structure	Content (%)	substance and ratio	Repeating structural unit, ratio	Weight average molecular weight		Water/ solvent
82	(2-1-2)/(B1-2) = 9/1	_	2%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-5)/ (F-1) = 9/1	6/4
83	(2-1-2)/(B1-3) = 9/1	_	2%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/ (F-2) = 9/1	6/4
84	(2-1-2)/(2-1-6)/(B1-1) = 3.5/3.5/3	_	2%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-5)/ (F-2) = 9/1	6/4
85	(2-1-2)/(2-1-6)/(B1-1) = 2.5/2.5/5	_	2%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)/ (F-2) = 9/1	6/4
86	(2-1-2)/(2-1-6)/(B1-1) = 1.5/1.5/7	_	2%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/ (F-1) = 9/1	6/4
87	(2-1-3)/(B-1) = 9/1		2%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-5)/ (F-2) = 9/1	6/4
88	(2-1-4)/(B1-1) = 9/1	_	2%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)/ (F-1) = 9/1	6/4
89	(2-1-5)/(B1-1) = 9/1	_	2%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/ (F-2) = 9/1	6/4
90	(2-1-6)/(B1-1) = 9/1		2%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)/ (F-1) = 9/1	6/4
91	(2-1-7)/(B1-1) = 9/1	_	2%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-5)/ (F-1) = 9/1	6/4
92	(2-1-1)/(2-1-4)/(B1-4) = 4.5/4.5/1	_	0.1%		(B1-5)	57000	(E-6)/ (F-1) = 9/1	6/4
93	(2-1-3)/(2-1-4)/(B1-5) = 4.5/4.5/1	_	1%	CTM-1/CTM-5 = 5/5	(B1-6)	57000	(E-6)/ (F-8) = 9/1	6/4
94	(2-1-5)/(2-1-6)/(B1-6) = 4.5/4.5/1	_	5%	CTM-1/CTM-5 = 3/7	(B1-7)	57000	(E-3)/ (F-14) = 9/1	6/4
95	(2-1-7)/(2-1-2)/(B1-7) = 4.5/4.5/1	_	2%	CTM-2/CTM-4 = 5/5	(B1-8)	57000	(F-14) = 9/1 (E-2)/ (F-33) = 9/1	6/4
96	(2-1-2)/(2-1-5)/(B1-8) = 4.5/4.5/1	_	2%	5/5 CTM-3/CTM-8 = 5/5	(B1-9)	57000	(F-33) = 9/1 (E-2)/ (F-18) = 9/1	6/4

TABLE 9

							Kind and ratio	of solvent
	Polyester havin	ng siloxane bond		Charge transporting	Binder	resin	Hydrophobic solvent/hydro-	
Example	Repeating structural unit, ratio	Terminal structure	Content (%)	substance and ratio	Repeating structural unit, ratio	Weight average molecular weight		Water/ solvent
97	(3-1-1)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-5)/ (F-1) = 9/1	6/4
98	(3-1-2)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/(F-2) = 9/1	6/4
99	(3-1-2)/(B2-2) = 9/1	(3-3-4)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)	6/4
100	(3-1-2)/(B2-3) = 9/1	(3-3-2) · (3-5)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)	6/4
101	(3-1-2)/(3-1-9)/(B2-1) = 3.5/3.5/3	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)	6/4
102	(3-1-2)(3-1-9)/(B2-1) = 2.5/2.5/5	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-5)/ (F-1) = 9/1	6/4
103	(3-1-2)/(3-1-9)/(B2-1) = 1.5/1.5/7	(3-3-2)	0.5%	$ \begin{array}{r} \text{CTM-1/CTM-7} = \\ 5/5 \end{array} $	(B1-1)	57000	(E-4)	6/4
104	(3-1-2)/(3-1-16)/(B2-1) = 1.5/1.5/7	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-5)/ (F-1) = 9/1	6/4
105	(3-1-3)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)	6/4
106	(3-1-4)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/(F-2) = 9/1	6/4
107	(3-1-5)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-5)	6/4
108	(3-1-6)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)/ (F-2) = 9/1	6/4
109	(3-1-7)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)/ (F-2) = 9/1	6/4
110	(3-1-9)/(B2-1) = 5/5	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-5)	6/4

TABLE 9-continued

							Kind and ratio	of solvent
	Polyester hav	ving siloxane bond		Charge _transporting	Binder	resin	Hydrophobic solvent/hydro-	
Example	Repeating structural unit, ratio	Terminal structure	Content (%)	substance and ratio	Repeating structural unit, ratio	Weight average molecular weight		Water/ solvent
111	(3-1-11)/(B2-1) = 7/3	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)/ (F-1) = 9/1	6/4
112	(3-1-14)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1) = 9/1 (E-1)	6/4
113	(3-1-16)/(B2-1) = 6/4	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-5)	6/4
114	(3-1-18)/(B2-1) = 8/2	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)/ (F-2) = 9/1	6/4
115	(3-1-21)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-5)	6/4
116	(3-1-2)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1	(B1-1)	57000	(E-4)	6/4
117	(3-1-2)/(B2-1) = 9/1 (3-1-2)/(B2-1) = 9/1	(3-3-2)	0.5%		(B1-1) (B1-1)	57000	(E-1)	6/4
118	(3-1-2)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 3/7	(B1-1)	57000	(E-5)/(F-2) = 9/1	6/4
119	(3-1-2)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-7	(B1-1)	57000	(E-1)	6/4
120	(3-1-2)/(B2-1) = 9/1	(3-3-2)	0.5%		(B1-1)	14000	(E-4)	6/4
121	(3-1-2)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-1)	120000	(E-5)/(F-2) = 9/1	6/4
122	(3-1-2)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-2)	55000	(E-4)	6/4
123	(3-1-2)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-3)	53000	(E-1)/ (F-1) = 9/1	6/4
124	(3-1-2)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B1-2)/(B1-3) = 3/7	55000	(F-1) = 9/1 (E-5)/ (F-2) = 9/1	6/4
125	(3-1-2)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5		55000	(E-4)/ (F-1) = 9/1	6/4
126	(3-1-2)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5		55000	(E-4)	6/4
127	(3-1-2)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5		120000	(E-1)/ (F-1) = 9/1	6/4
128	(3-1-2)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B2-2)	120000	(F-1) = 9/1 (E-1)/ (F-2) = 9/1	6/4
129	(3-1-2)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5	(B2-1)/(B2-2) = 7/3	120000	(E-4)	6/4
130	(3-1-2)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 = 5/5		57000	(E-4)/ (F-1) = 9/1	6/4
131	(3-1-2)/(B2-1) = 9/1	(3-3-2)	0.5%	CTM-1/CTM-7 =	(B1-1)	57000	(E-1)/	6/4
132	(3-1-2)/(B2-1) = 9/1	(3-3-2)	0.5%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-1) = 9/1 (F-1)	6/4
133	(3-1-2)/(B2-4) = 9	(3-3-1)	0.1%	5/5 CTM-1/CTM-5 =	· / / /	57000	(E-6)/	6/4
134	(3-1-2)/(B2-5) = 9	(3-3-2) · (3-4)	1%	3/7 CTM-1/CTM-5 =		57000	(F-6) = 9/1 (E-6)/	6/4
135	(3-1-2)/(B2-6) = 9	(3-3-2)	5%	5/5 CTM-1/CTM-5 =	\ /\ /	57000	(F-35) = 9/1 (E-3)/	6/4
136	(3-1-2)/(B2-1) = 9/1	(3-3-2)	1%	7/3 CTM-8/CTM-9 = 5/5	7/3 (B2-4)/(B2-6) = 5/5	57000	(F-23) = 9/1 (E-6)/ (F-29) = 9/1	6/4

TABLE 10

						Kind and ratio	of solvent
	Polystyrene having sile	oxane bond	Charge _transporting	Binder	resin	Hydrophobic _solvent/hydro-	
Example	Repeating structural unit, ratio	Content (%)	substance and ratio	Repeating structural unit, ratio	Weight average molecular weight		Water/ solvent
137	(4-1-1)/(4-2) = 2/8	1%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)/ (F-1) = 9/1	6/4
138	(4-1-2)/(4-2) = 2/8	1%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/ (F-1) = 9/1	6/4
139	(4-1-3)/(4-2) = 1/9	1%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)	6/4
140	(4-1-3)/(4-2) = 2/8	1%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-5)/ (F-2) = 9/1	6/4

TABLE 10-continued

						Kind and ratio	of solvent
	Polystyrene having sil	oxane bond	Charge transporting	Binder	resin	Hydrophobic _solvent/hydro-	
Example	Repeating structural unit, ratio	Content (%)	substance and ratio	Repeating structural unit, ratio	Weight average molecular weight		Water/ solvent
141	(4-1-3)/(4-2) = 3/7	1%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-5)	6/4
142	(4-1-4)/(4-2) = 2/8	1%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-5)/ (F-2) = 9/1	6/4
143	(4-1-5)/(4-2) = 2/8	1%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)	6/4
144	(4-1-3)/(4-2) = 2/8	1%	CTM-1	(B1-1)	57000	(E-4)/ (F-1) = 9/1	6/4
145	(4-1-3)/(4-2) = 2/8	1%	CTM-1/CTM-7 = 7/3	(B1-1)	57000	(E-4)	6/4
146	(4-1-3)/(4-2) = 2/8	1%	CTM-1/CTM-7 = 3/7	(B1-1)	57000	(E-5)	6/4
147	(4-1-3)/(4-2) = 2/8	1%	CTM-7	(B1-1)	57000	(E-1)/	6/4
148	(4-1-3)/(4-2) = 2/8	1%	CTM-1/CTM-7 =	(B1-1)	14000	(F-2) = 9/1 (E-1)/	6/4
149	(4-1-3)/(4-2) = 2/8	1%	5/5 CTM-1/CTM-7 =	(B1-1)	120000	(F-2) = 9/1 (E-1)	6/4
150	(4-1-3)/(4-2) = 2/8	1%	5/5 CTM-1/CTM-7 =	(B1-2)	55000	(E-4)/	6/4
151	(4-1-3)/(4-2) = 2/8	1%	5/5 CTM-1/CTM-7 =	(B1-3)	53000	(F-2) = 9/1 (E-5)/	6/4
152	(4-1-3)/(4-2) = 2/8	1%	5/5 CTM-1/CTM-7 =		55000	(F-1) = 9/1 (E-4)	6/4
153	(4-1-3)/(4-2) = 2/8	1%	5/5 CTM-1/CTM-7 =		55000	(E-1)/	6/4
154	(4-1-3)/(4-2) = 2/8	1%		5/5 (B1-2)/(B1-3) =	55000	(F-1) = 9/1 (E-4)	6/4
155	(4-1-3)/(4-2) = 2/8	1%	5/5 CTM-1/CTM-7 =	7/3 (B2-1)	120000	(E-5)/	6/4
156	(4-1-3)/(4-2) = 2/8	1%	5/5 CTM-1/CTM-7 =	(B2-2)	120000	(F-1) = 9/1 (E-1)/	6/4
157	(4-1-3)/(4-2) = 2/8	1%	5/5 CTM-1/CTM-7 =		120000	(F-1) = 9/1 (E-4)	6/4
158	(4-1-3)/(4-2) = 2/8	1%	5/5 CTM-1/CTM-7 =	7/3 (B1-1)	57000	(E-4)/	6/4
159	(4-1-3)/(4-2) = 2/8	1%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-2) = 9/1 (E-5)	6/4
160	(4-1-3)/(4-2) = 2/8	1%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-1)	6/4
161	(4-1-3)/(4-2) = 2/8	0.5%	5/5 CTM-1/CTM-5 =	(B1-4)	57000	(E-6)/	6/4
162	(4-1-3)/(4-2) = 2/8	3%	3/7 CTM-1/CTM-5 =	(B1-5)	57000	(F-17) = 9/1 (E-2)/	6/4
163	(4-1-3)/(4-2) = 2/8	10%	5/5 CTM-1/CTM-5 =	,	57000	(F-30) = 9/1 (E-2)/	6/4
164		0.5%	7/3 CTM-2/CTM-3 =		57000	(E-2)/ (F-35) = 9/1 (E-6)/	6/4
	(4-1-3)/(4-2) = 2/8		5/5	,		(F-26) = 9/1	
165	(4-1-3)/(4-2) = 2/8	3%	CTM-6/CTM-8 = 5/5	(B2-5)	57000	(E-6)/ (F-15) = 9/1	6/4

TABLE 11

						Kind and ratio of	f solvent
	Compound by form	represented iula (5)	Charge _transporting	Binder	resin	Hydrophobic solvent/hydro-	
Example	Formula	Content (%)	substance and ratio	Repeating structural unit, ratio	Weight average molecular weight		Water/ solvent
166	(5-1)	1%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)/ (F-2) = 9/1	6/4
167	(5-2)	10%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)/ (F-2) = 9/1	6/4
168	(5-3)	1%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/ (F-2) = 9/1	6/4
169	(5-4)	10%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/ (F-2) = 9/1	6/4

TABLE 11-continued

						Kind and ratio	of solvent
		l represented mula (5)	l Charge _transporting	Binder	resin	Hydrophobic solvent/hydro-	
Example	Formula	Content (%)	substance and ratio	Repeating structural unit, ratio	Weight average molecular weight		Water/ solvent
170	(5-5)	2%	CTM-1/CTM-7 =	(B1-1)	57000	(E-5)	6/4
171	(5-6)	2%	5/5 CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)/ (F-1) = 9/1	6/4
172	(5-7)	2%	CTM-1/CTM-7 =	(B1-1)	57000	(E-1)/	6/4
173	(5-2)	2%	5/5 CTM-1	(B1-1)	57000	(F-1) = 9/1 (E-5)/	6/4
174	(5-2)	2%	CTM-1/CTM-7 =	(B1-1)	57000	(F-2) = 9/1 (E-1)/	6/4
175	(5-2)	2%		(B1-1)	57000	(F-2) = 9/1 (E-5)/	6/4
176	(5-2)	2%	3/7 CTM-7	(B1-1)	57000	(F-2) = 9/1 (E-1)/	6/4
177	(5-2)	2%	CTM-1/CTM-7 =	(B1-1)	14000	(F-1) = 9/1 (E-5)	6/4
178	(5-2)	2%	5/5 CTM-1/CTM-7 =	(B1-1)	120000	(E-4)/	6/4
179	(5-2)	2%	5/5 CTM-1/CTM-7 =	(B1-2)	55000	(F-1) = 9/1 (E-1)/	6/4
180	(5-2)	2%	5/5 CTM-1/CTM-7 =	(B1-3)	53000	(F-1) = 9/1 (E-5)/	6/4
181	(5-2)	2%	5/5 CTM-1/CTM-7 =		55000	(F-2) = 9/1 (E-5)/	6/4
182	(5-2)	2%	5/5 CTM-1/CTM-7 =		55000	(F-1) = 9/1 (E-1)	6/4
183	(5-2)	2%	5/5 CTM-1/CTM-7 =		55000	(E-4)/	6/4
184	(5-2)	2%	5/5 CTM-1/CTM-7 =	7/3 (B2-1)	120000	(F-1) = 9/1 (E-5)/	6/4
185	(5-2)	2%	5/5 CTM-1/CTM-7 =	(B2-2)	120000	(F-1) = 9/1 (E-1)	6/4
186	(5-2)	2%	5/5 CTM-1/CTM-7 =	(B2-1)/(B2-2) = 7/3	120000	(E-4)/	6/4
187	(5-2)	2%	5/5 CTM-1/CTM-7 = 5/5		57000	(F-2) = 9/1 (E-5)/	6/4
188	(5-2)	2%	CTM-1/CTM-7 =	(B1-1)	57000	(F-1) = 9/1 (E-4)	6/4
189	(5-2)	2%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-1)	6/4
190	(5-2)	0.5%	5/5 CTM-1/CTM-5 =	(B1-4)	57000	(E-6)/	6/4
191	(5-2)	2%	7/3 CTM-1/CTM-5 =	(B1-5)	57000	(F-4) = 9/1 (E-3)/	6/4
192	(5-2)	5%	5/5 CTM-1/CTM-5 =	(B1-6)	57000	(F-19) = 9/1 (E-2)/	6/4
193	(5-2)	10%	3/7 CTM-2/CTM-3 =	(B1-7)	57000	(F-28) = 9/1 (E-2)/	6/4
194	(5-2)	1%	5/5 CTM-4/CTM-6 =	(B1-8)	57000	(F-31) = 9/1 (E-6)/	6/4
195	(5-2)	5%	5/5 CTM-8/CTM-9 =	, ,	57000	(F-12) = 9/1 (E-2)/	6/4
170	(Z)	370	5/5	(-1-2)	2,000	(F-13) = 9/1	V/T

TABLE 12

						Kind and ratio	of solvent
		represented nula (6)	Charge _transporting	Binder	resin	Hydrophobic _solvent/hydro-	
Example	Formula	Content (%)	substance and ratio	Repeating structural unit, ratio	Weight average molecular weight		Water/ solvent
196	(6-1)	1%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/ (F-2) = 9/1	6/4
197	(6-2)	10%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/ (F-2) = 9/1	6/4
198	(6-3)	1%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)/ (F-1) = 9/1	6/4

TABLE 12-continued

						Kind and ratio	of solvent
		represented nula (6)	Charge _transporting	Binder	resin	Hydrophobic _solvent/hydro-	
Example	Formula	Content (%)	substance and ratio	Repeating structural unit, ratio	Weight average molecular weight		Water/ solvent
199	(6-4)	10%	CTM-1/CTM-7 =	(B1-1)	57000	(E-4)/	6/4
200	(6-5)	3%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-1) = 9/1 (E-1)/	6/4
201	(6-6)	3%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-1) = 9/1 (E-4)	6/4
202	(6-7)	3%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(E-4)/	6/4
203	(6-4)	3%	5/5 CTM-1	(B1-1)	57000	(F-2) = 9/1 (E-1)/	6/4
204	(6-4)	3%	CTM-1/CTM-7 =	(B1-1)	57000	(F-1) = 9/1 (E-5)/	6/4
205	(6-4)	3%	7/3 CTM-1/CTM-7 =	(B1-1)	57000	(F-2) = 9/1 (E-1)/	6/4
206	(6-4)	3%	3/7 CTM-7	(B1-1)	57000	(F-2) = 9/1 (E-4)/	6/4
207	(6-4)	3%	CTM-1/CTM-7 =	(B1-1)	14000	(F-1) = 9/1 (E-5)/	6/4
208	(6-4)	3%	5/5 CTM-1/CTM-7 =	(B1-1)	120000	(F-2) = 9/1 (E-1)/	6/4
209	(6-4)	3%	5/5 CTM-1/CTM-7 =	(B1-2)	55000	(F-1) = 9/1 (E-4)/	6/4
210	(6-4)	3%	5/5 CTM-1/CTM-7 =	(B1-3)	53000	(F-2) = 9/1 (E-4)/	6/4
211	(6-4)	3%	5/5 CTM-1/CTM-7 =	(B1-2)/(B1-3) =	55000	(F-1) = 9/1 (E-5)/	6/4
212	(6-4)	3%	5/5 CTM-1/CTM-7 =	3/7 (B1-2)/(B1-3) =	55000	(F-2) = 9/1 (E-5)/	6/4
213	(6-4)	3%	5/5 CTM-1/CTM-7 =	5/5 (B1-2)/(B1-3) =	55000	(F-2) = 9/1 (E-1)	6/4
214	(6-4)	3%	5/5 CTM-1/CTM-7 =	7/3 (B2-1)	120000	(E-5)/	6/4
215	(6-4)	3%	5/5 CTM-1/CTM-7 =	(B2-2)	120000	(F-2) = 9/1 (E-5)	6/4
216	(6-4)	3%	5/5 CTM-1/CTM-7 =	(B2-1)/(B2-2) =	120000	(E-5)/	6/4
217	(6-4)	3%	5/5 CTM-1/CTM-7 =	7/3 (B1-1)	57000	(F-1) = 9/1 (E-1)/	6/4
218	(6-4)	3%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-1) = 9/1 (E-4)/	6/4
219	(6-4)	3%	5/5 CTM-1/CTM-7 =		57000	(F-2) = 9/1 (F-1)	6/4
220	(6-4)	1%	5/5 CTM-1/CTM-5 =		57000	(E-3)/	6/4
	, ,		7/3	5/5		(F-22) = 9/1	
221	(6-4)	3%	CTM-1/CTM-5 = 5/5	5/5	57000	(E-6)/ (F-27) = 9/1	6/4
222	(6-4)	10%	CTM-1/CTM-5 = 3/7	(B2-4)/(B2-5) = 5/5	57000	(E-2)/ (F-34) = 9/1	6/4
223	(6-4)	5%	CTM-4/CTM-8 = 5/5	(B1-4)/(B1-8) = 5/5	57000	(E-3)/ (F-24) = 9/1	6/4
224	(6-4)	5%	CTM-3/CTM-9 = 5/5		57000	(E-6)/ (F-9) = 9/1	6/4

TABLE 13

						Kind and ratio of	of solvent
	Compound :		Charge _transporting	Binder	resin	Hydrophobic _solvent/hydro-	
Example	Formula	Content (%)	substance and ratio	Repeating structural unit, ratio	Weight average molecular weight		Water/ solvent
225	(7-1-1)	1%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/ (F-1) = 9/1	6/4
226	(7-1-2)	10%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/ (F-1) = 9/1	6/4
227	(7-1-3)	1%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)/ (F-1) = 9/1	6/4

			TAB	LE 13-continued			
						Kind and ratio	of solvent
	Compound by forn		l Charge _transporting	Binder	resin	Hydrophobic _solvent/hydro-	
Example	Formula	Content (%)	substance and ratio	Repeating structural unit, ratio	Weight average molecular weight		Water/ solvent
228	(7-1-4)	10%	CTM-1/CTM-7 =	(B1-1)	57000	(E-4)/	6/4
229	(7-1-5)	3%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-1) = 9/1 (E-1)	6/4
230	(7-1-6)	3%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(E-4)	6/4
231	(7-1-7)	3%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(E-5)/	6/4
232	(7-1-5)	3%	5/5 CTM-1	(B1-1)	57000	(F-1) = 9/1 (E-4)/	6/4
233	(7-1-6)	3%	CTM-1/CTM-7 =	(B1-1)	57000	(F-2) = 9/1 (E-4)	6/4
234	(7-1-7)	3%	7/3 CTM-1/CTM-7 =	(B1-1)	57000	(E-5)/	6/4
235	(7-1-8)	3%	3/7 CTM-7	(B1-1)	57000	(F-2) = 9/1 (E-1)/	6/4
236	(7-1-9)	3%	CTM-1/CTM-7 =	(B1-1)	14000	(F-1) = 9/1 (E-4)	6/4
237	(7-1-10)	3%	5/5 CTM-1/CTM-7 =	(B1-1)	120000	(E-5)/	6/4
238	(7-1-11)	3%	5/5 CTM-1/CTM-7 =	(B1-2)	55000	(F-1) = 9/1 (E-5)	6/4
239	(7-1-12)	3%	5/5 CTM-1/CTM-7 =	(B1-3)	53000	(E-1)/	6/4
240	(7-1-13)	3%	5/5 CTM-1/CTM-7 =	(B1-2)/(B1-3) =	55000	(F-2) = 9/1 (E-4)	6/4
241	(7-1-14)	3%	5/5 CTM-1/CTM-7 =	3/7 (B1-2)/(B1-3) =	55000	(E-4)/	6/4
242	(7-1-15)	3%	5/5 CTM-1/CTM-7 =	5/5 (B1-2)/(B1-3) =	55000	(F-1) = 9/1 (E-5)/	6/4
243	(7-1-16)	3%	5/5 CTM-1/CTM-7 =	7/3 (B2-1)	120000	(F-1) = 9/1 (E-1)	6/4
244	(7-1-17)	3%	5/5 CTM-1/CTM-7 =	(B2-2)	120000	(E-5)/	6/4
245	(7-1-18)	3%	5/5 CTM-1/CTM-7 =	(B2-1)/(B2-2) =	120000	(F-2) = 9/1 (E-1)	6/4
246	(7-1-19)	3%	5/5 CTM-1/CTM-7 =	7/3 (B1-1)	57000	(E-5)/	6!4
247	(7-1-20)	3%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-1) = 9/1 (E-4)/	6/4
248	(7-1-28)	3%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-2) = 9/1 (F-1)	6/4
249	(7-1-8)	1%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(E-1)/	6/4
250	(7-1-9)	10%	5/5 CTM-1/CTM-7 =		57000	(F-1) = 9/1 (E-1)/	6/4
251	(7-1-10)	1%	5/5 CTM-1/CTM-7 =		57000	(F-1) = 9/1 (E-4)/	6/4
252	(7-1-11)	10%	5/5 CTM-1/CTM-7 =		57000	(F-1) = 9/1 (E-4)/	6/4
253	(7-1-12)	3%	5/5 CTM-1/CTM-7 =	,	57000	(F-1) = 9/1 (E-1)	6/4
254	(7-1-13)	3%	5/5 CTM-1/CTM-7 =	,	57000	(E-5)/	6/4
255	(7-1-10)	3%	5/5 CTM-1	(B1-1)	57000	(F-1) = 9/1 (E-1)/	6/4
				,		(F-1) = 9/1	
256	(7-1-10)	3%	CTM-1/CTM-7 = 7/3	,	57000	(E-4)/ (F-2) = 9/1	6/4
257	(7-1-10)	3%	CTM-1/CTM-7 = 3/7	(B1-1)	57000	(E-1)	6/4
258 259	(7-1-10) (7-1-10)	3% 3%	CTM-7 CTM-1/CTM-7 =	(B1-1) (B1-1)	57000 14000	(E-1) (E-5)/	6/4 6/4
260	(7-1-10)	3%	5/5 CTM-1/CTM-7 =	(B1-1)	120000	(F-2) = 9/1 (E-5)/	6/4
261	(7-1-10)	3%	5/5 CTM-1/CTM-7 =	, ,	55000	(F-1) = 9/1 (E-4)/	6/4
			5/5	,	53000	(F-1) = 9/1 (E-1)/	6/4
262	(7-1-10)	3%	CTM-1/CTM-7 = 5/5	,		(F-2) = 9/1	
263	(7-1-10)	3%	CTM-1/CTM-7 = 5/5	(B1-2)/(B1-3) = 3/7	55000	(E-5)/ (F-2) = 9/1	6/4

TABLE 13-continued

						Kind and ratio	of solven
	1	l represented mula (7)	l Charge _transporting	Binder	resin	Hydrophobic solvent/hydro-	
Example	Formula	Content (%)	substance and ratio	Repeating structural unit, ratio	Weight average molecular weight		Water/ solvent
264	(7-1-10)	3%	CTM-1/CTM-7 = 5/5	(B1-2)/(B1-3) = 5/5	55000	(E-5)	6/4
265	(7-1-10)	3%	CTM-1/CTM-7 = 5/5		55000	(E-4)	6/4
266	(7-1-10)	3%	CTM-1/CTM-7 = 5/5		120000	(E-1)/ (F-2) = 9/1	6/4
267	(7-1-10)	3%	CTM-1/CTM-7 = 5/5	(B2-2)	120000	(E-5)/ (F-1) = 9/1	6/4
268	(7-1-10)	3%	CTM-1/CTM-7 = 5/5	(B2-1)/(B2-2) = 7/3	120000	(E-4)/ (F-2) = 9/1	6/4
269	(7-1-10)	3%	CTM-1/CTM-7 = 5/5		57000	(E-5)	6/4
270	(7-1-10)	3%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)	6/4
271	(7-1-10)	3%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(F-1)	6/4
272	(7-1-14)	1%	5/5 CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-1)/ (F-1) = 9/1	6/4
273	(7-1-15)	10%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(F-1) = 9/1 (E-1)/ (F-1) = 9/1	6/4
274	(7-1-16)	1%	CTM-1/CTM-7 = 5/5	(B1-1)	57000	(E-4)/	6/4
275	(7-1-17)	10%	CTM-1/CTM-7 =	(B1-1)	57000	(F-1) = 9/1 (E-4)/	6/4
276	(7-1-18)	3%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-1) = 9/1 (E-1)/	6/4
277	(7-1-19)	3%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-2) = 9/1 (E-5)	6/4
278	(7-1-20)	3%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(E-1)/	6/4
279	(7-1-21)	3%	5/5 CTM-1/CTM-7 =	(B1-1)	57000	(F-2) = 9/1 (E-5)/	6/4
280	(7-1-16)	3%	5/5 CTM-1	(B1-1)	57000	(F-1) = 9/1 (E-4)/	6/4
281	(7-1-16)	3%	CTM-1/CTM-7 =	(B1-1)	57000	(F-1) = 9/1 (E-5)	6/4
282	(7-1-16)	3%	7/3 CTM-1/CTM-7 =	(B1-1)	57000	(E-1)/	6/4
283	(7-1-16)	3%	3/7 CTM-7	(B1-1)	57000	(F-1) = 9/1 (E-5)/	6/4
284	(7-1-16)	3%	CTM-1/CTM-7 =	(B1-1)	14000	(F-2) = 9/1 (E-5)/	6/4
285	(7-1-16)	3%	5/5 CTM-1/CTM-7 =	(B1-1)	120000	(F-2) = 9/1 (E-4)	6/4
286	(7-1-16)	3%	5/5 CTM-1/CTM-7 =	(B1-2)	55000	(E-5)/	6/4
287	(7-1-16)	3%	5/5 CTM-1/CTM-7 =	(B1-3)	53000	(F-1) = 9/1 (E-4)/	6/4
288	(7-1-16)	3%	5/5 CTM-1/CTM-7 =	(B1-2)/(B1-3) =	55000	(F-2) = 9/1 (E-1)	6/4
289	(7-1-16)	3%	5/5 CTM-1/CTM-7 =	3/7 (B1-2)/(B1-3) =	55000	(E-4)/	6/4
290	(7-1-16)	3%	5/5 CTM-1/CTM-7 =	5/5 (B1-2)/(B1-3) =	55000	(F-2) = 9/1 (E-1)	6/4
291	(7-1-16)	3%	5/5 CTM-1/CTM-7 =	7/3 (B2-1)	120000	(E-5)	6/4
292	(7-1-16)	3%	5/5 CTM-1/CTM-7 =	(B2-2)	120000	(E-1)	6/4
293	(7-1-16)	3%	5/5 CTM-1/CTM-7 =		120000	(E-4)	6/4
294	(7-1-16)	3%	5/5 CTM-1/CTM-7 =	7/3	57000	(E-4)	6/4
295	(7-1-16)	3%	5/5 CTM-1/CTM-7 =		57000	(E-5)/	6/4
	,		5/5	, ,		(F-1) = 9/1	
296	(7-1-16)	3%	CTM-1/CTM-7 = 5/5	(D1-1)	57000	(F-1)	6/4

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TABLE 14-continued

		TABLE	14			TABLE 14-continued					
		Evaluation of	f solution stability					Evaluation of	solution stability		
	Immediately af	ter preparation	Leaving for 2 w	eeks and stirring	5		Immediately af	ter preparation	Leaving for 2 w	eeks and stirring	
Exam- ple	Visual observation	Average particle diameter	Visual observation	Average particle diameter		Exam- ple	Visual observation		Visual observation	Average particle diameter	
1	Uniform and semi-	2.2 µm	Uniform and semi-	2.5 μm	10	31	Uniform and semi-	3.2 µm	Uniform and semi-	3.4 µm	
2	transparent Uniform and transparent	1.1 µm	transparent Uniform and transparent	1.3 μm		32	transparent Uniform and semi-	3.2 µm	transparent Uniform and semi-	3.5 µm	
3	Uniform and transparent	0.9 μm	Uniform and transparent	1.0 μm		33	transparent Uniform and	3.7 µm	transparent Uniform and	3.9 µm	
4	Uniform and semi- transparent	2.1 µm	Uniform and semi- transparent	2.4 μm	15	34	semi- transparent Uniform and	3.4 µm	semi- transparent Uniform and	3.7 µm	
5	Uniform and transparent	0.8 μm	Uniform and transparent	0.9 μm		2.5	semi- transparent	·	semi- transparent	·	
6 7	Uniform and transparent Uniform and	0.8 μm 1.2 μm	Uniform and transparent Uniform and	0.9 μm 1.4 μm	20	35	Uniform and semi- transparent	2.5 µm	Uniform and semi- transparent	2.8 µm	
8	transparent Uniform and	2.4 μm	transparent Uniform and	2.7 μm		36	Uniform and transparent	0.8 μm	Uniform and transparent	0.9 μm	
9	semi- transparent Uniform and	1.2 µm	semi- transparent Uniform and	1.3 μm		37 38	Uniform and transparent Uniform and	1.0 µm 0.9 µm	Uniform and transparent Uniform and	1.1 μm 1.0 μm	
10	transparent Uniform and	1.0 µm	transparent Uniform and	1.2 μm	25	39	transparent Uniform and	2.4 μm	transparent Uniform and	2.7 μm	
11	transparent Uniform and transparent	1.4 µm	transparent Uniform and transparent	1.5 μm		40	semi- transparent Uniform and	1.2 μm	semi- transparent Uniform and	1.3 μm	
12	Uniform and transparent	1.6 µm	Uniform and transparent	1.8 μm	30	41	transparent Uniform and	1.6 μm	transparent Uniform and	1.8 µm	
13 14	Uniform and transparent Uniform and	1.4 µm 2.0 µm	Uniform and transparent Uniform and	1.5 μm 2.3 μm		42	transparent Uniform and transparent	1.5 μm	transparent Uniform and transparent	1.6 μm	
14	semi- transparent	2.0 μπ	semi- transparent	2.5 μπ		43	Uniform and semi-	2.8 µm	Uniform and semi-	3.0 µm	
15 16	Uniform and transparent Uniform and	1.2 µm 1.0 µm	Uniform and transparent Uniform and	1.4 μm 1.3 μm	35	44	transparent Uniform and transparent	1 μm	transparent Uniform and transparent	1.1 μm	
17	transparent Uniform and	1.0 µm	transparent Uniform and	1.3 μm		45	Uniform and semi-	2.3 µm	Uniform and semi-	2.5 μm	
18	transparent Uniform and	0.9 µm	transparent Uniform and transparent	1.0 μm	40	46	transparent Uniform and	1.2 μm	transparent Uniform and	1.3 μm	
19	transparent Uniform and semi-	2.1 µm	Uniform and semi-	2.3 µm		47	transparent Uniform and semi-	2.7 µm	transparent Uniform and semi-	2.8 µm	
20	transparent Uniform and	1.5 µm	transparent Uniform and transparent	1.7 µm		48	transparent Uniform and transparent	1.5 µm	transparent Uniform and transparent	1.6 µm	
21	transparent Uniform and transparent	1.7 µm	Uniform and transparent	1.8 µm	45	49	Uniform and transparent	1.6 µm	Uniform and transparent	1.8 μm	
22	Uniform and semi-	2.4 µm	Uniform and semi-	2.6 µm		50	Uniform and semi-	2.6 µm	Uniform and semi-	2.8 µm	
23	transparent Uniform and transparent	1.7 µm	transparent Uniform and transparent	1.8 µm	50	51	transparent Uniform and transparent	1.3 µm	transparent Uniform and transparent	1.4 μm	
24	Uniform blue white	4.1 μm	Uniform blue white	4.5 μm		52	Uniform and transparent	1.1 μm	Uniform and transparent	1.2 μm	
25	Uniform and semi- transparent	3.7 µm	Uniform and semi- transparent	3.9 µm		53 54	Uniform and transparent Uniform and	0.9 μm 2.4 μm	Uniform and transparent Uniform and	1.0 μm 2.5 μm	
26	Uniform and semi-	3.4 µm	Uniform and semi-	3.7 µm	55		semi- transparent		semi- transparent		
27	transparent Uniform and semi-	3.1 µm	transparent Uniform and semi-	3.4 µm		55 56	Uniform and transparent Uniform and	1.6 µm 2.2 µm	Uniform and transparent Uniform and	1.8 μm 2.4 μm	
28	transparent Uniform and semi-	3.2 μm	transparent Uniform and semi-	3.4 µm	60	57	semi- transparent Uniform and		semi- transparent Uniform and	·	
29	transparent Uniform and	3.1 µm	transparent Uniform and	3.3 µm			semi- transparent	2.7 μm	semi- transparent	3.0 µm	
30	semi- transparent Uniform and	3.2 µm	semi- transparent Uniform and	3.4 µm		58 59	Uniform and transparent Uniform and	1.2 µm 2.4 µm	Uniform and transparent Uniform and	1.4 μm 2.7 μm	
23	semi- transparent	3.2 pm	semi- transparent	puii	65		semi- transparent	2 part	semi- transparent	2., pm1	

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TABLE 14-continued

		ABLE 14-c						ABLE 14-c		
		Evaluation of	solution stability					Evaluation of	solution stability	
	Immediately aft	er preparation	Leaving for 2 week	s and stirring	5		Immediately at	ter preparation	Leaving for 2 w	eeks and stirring
Exam- ple	Visual observation	Average particle diameter	Visual observation	Average particle diameter		Exam- ple	Visual observation	Average particle diameter	Visual observation	Average particle diameter
60	Uniform and semi-	2.6 μm	Uniform and semi-	2.8 µm	10	91	Uniform and transparent	1.2 μm	Uniform and transparent	1.4 µm
61	transparent Uniform and semi-	2.3 µm	transparent Uniform and semi-	2.5 µm		92	Uniform and semi- transparent	3.5 μm	Uniform and semi- transparent	3.9 μm
62	transparent Uniform and semi-	2.2 μm	transparent Uniform and semi-	2.4 μm	15	93	Uniform and semi- transparent	3.2 µm	Uniform and semi- transparent	3.5 μm
63	transparent Uniform and transparent	1.1 µm	transparent Uniform and transparent	1.3 μm	13	94	Uniform and semi- transparent	3.4 µm	Uniform and semi- transparent	3.6 µm
64	Uniform and transparent	1.7 µm	Uniform and transparent	1.9 μm		95	Uniform and semi-	3.2 μm	Uniform and semi-	3.5 µm
65	Uniform and transparent	1.5 μm	Uniform and transparent	1.6 μm	20	96	transparent Uniform and	3.2 μm	transparent Uniform and	3.5 µm
66	Uniform and semi- transparent	2.4 μm	Uniform and semi- transparent	2.6 μm		97	semi- transparent Uniform and	1.2 μm	semi- transparent Uniform and	1.4 μm
67	Uniform and transparent	,	Uniform and transparent	1.3 µm		98	transparent Uniform and	1.5 μm	transparent Uniform and	1.6 μm
68	Uniform and semi- transparent	2.1 μm	Uniform and semi- transparent	2.3 μm	25	99	transparent Uniform and semi-	2.3 μm	transparent Uniform and semi-	2.5 μm
69	Uniform and semi-	2.6 μm	Uniform and semi-	2.8 μm		100	transparent Uniform and semi-	2.5 μm	transparent Uniform and semi-	2.6 μm
70	transparent Uniform and transparent		transparent Uniform and transparent	1.3 μm	30	101	transparent Uniform and	2.4 μm	transparent Uniform and	2.6 μm
71 72	Uniform and transparent Uniform and		Uniform and transparent Uniform and	1.1 μm 3.0 μm		102	semi- transparent Uniform and	1.7 µm	semi- transparent Uniform and	1.9 µm
, 2	semi- transparent	2.0 μm	semi- transparent	5.0 µп	35	103	transparent Uniform and	2.7 μm	transparent Uniform and	3.0 µm
73	Uniform and transparent		Uniform and transparent	1.9 μm	33	101	semi- transparent		semi- transparent	
74 75	Uniform and transparent Uniform and		Uniform and transparent Uniform and	1.5 µm 2.8 µm		104 105	Uniform and transparent Uniform and	1.3 μm 2.5 μm	Uniform and transparent Uniform and	1.5 μm 2.8 μm
,,,	semi- transparent	2.0 μm	semi- transparent	2.0 діп	40	103	semi- transparent	2.5 µm	semi- transparent	2.0 μm
76	Uniform and transparent		Uniform and transparent	1.3 μm		106	Uniform and transparent	0.8 μm	Uniform and transparent	0.9 μm
77	Uniform and semi- transparent	2.9 µm	Uniform and semi- transparent	3.1 μm		107	Uniform and semi- transparent	2.4 μm	Uniform and semi- transparent	2.6 µm
78	Uniform and transparent		Uniform and transparent	2.0 μm	45	108	Uniform and transparent	1.1 µm	Uniform and transparent	1.2 µm
79 80	Uniform blue white Uniform and		Uniform blue white Uniform and	4.6 µm 1.0 µm		109 110	Uniform and transparent Uniform and	1.0 μm 2.7 μm	Uniform and transparent Uniform and	1.0 μm 2.8 μm
81	transparent Uniform and	·	transparent Uniform and	1.0 μm 0.9 μm	50	110	semi- transparent	2.7 µm	semi- transparent	2.6 μπ
82	transparent Uniform and	•	transparent Uniform and	1.4 μm	50	111	Uniform and transparent	1.4 μm	Uniform and transparent	1.5 µm
83	transparent Uniform and	•	transparent Uniform and	1.3 μm		112	Uniform and semi-	2.8 µm	Uniform and semi-	3.0 µm
84	transparent Uniform and transparent	1.4 μm	transparent Uniform and transparent	1.6 μm	55	113	transparent Uniform and semi-	2.2 μm	transparent Uniform and semi-	2.3 µm
85	Uniform and transparent	1.3 μm	Uniform and transparent	1.3 μm		114	transparent Uniform and	1.7 µm	transparent Uniform and	1.9 µm
86	Uniform and transparent	1.7 µm	Uniform and transparent	1.8 µm		115	transparent Uniform and	2.4 μm	transparent Uniform and	2.7 μm
87	Uniform and transparent	·	Uniform and transparent	1.3 μm	60		semi- transparent		semi- transparent	
88	Uniform and transparent	·	Uniform and transparent	1.9 μm		116	Uniform and semi-	2.1 µm	Uniform and semi-	2.4 µm
89	Uniform and transparent	,	Uniform and transparent	1.5 μm	65	117	transparent Uniform and	2.7 µm	transparent Uniform and	2.9 µm
90	Uniform and transparent	0.8 µm	Uniform and transparent	0.9 μm	0.5		semi- transparent		semi- transparent	

TABLE 14-continued

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TABLE 14-continued

		Evaluation of	solution stability					Evaluation of	solution stability	
	Immediately at	fter preparation	Leaving for 2 w	eeks and stirring	5		Immediately a	ifter preparation	Leaving for 2 w	eeks and stirring
Exam- ple	Visual observation	Average particle diameter	Visual observation	Average particle diameter	,	Exam- ple	Visual observation	Average particle	Visual observation	Average particle diameter
118	Uniform and	0.9 µm	Uniform and	1.0 μm						
119	transparent Uniform and semi-	2.6 µm	transparent Uniform and semi-	2.8 µm	10	147 148	Uniform and transparent Uniform and	1.3 μm 0.8 μm	Uniform and transparent Uniform and	1.5 μm 0.9 μm
120	transparent Uniform and semi-	2.5 μm	transparent Uniform and semi-	2.8 µm		149	transparent Uniform and	2.1 μm	transparent Uniform and	2.3 μm
121	transparent Uniform and	1.2 µm	transparent Uniform and	1.4 μm	15		semi- transparent		semi- transparent	
122	transparent Uniform and semi-	2.2 µm	transparent Uniform and semi-	2.5 μm		150	Uniform and transparent	1.1 μm	Uniform and transparent	1.3 μm
123	transparent Uniform and transparent	1.5 μm	transparent Uniform and transparent	1.8 μm	20					
124	Uniform and	1.6 µm	Uniform and	1.9 µm				TABLE	15	
125	transparent Uniform and transparent	0.8 μm	transparent Uniform and transparent	0.9 μm				Evaluation o	of solution stability	y
126	Uniform and semi-	2.3 µm	Uniform and semi-	2.5 μm	25			mediately preparation		r 2 weeks and rring
127	transparent Uniform and transparent	1.2 µm	transparent Uniform and transparent	1.4 μm			Visual	Average particle		Average particle
128	Uniform and transparent	1.1 µm	Uniform and transparent	1.2 μm		Examp	le observation	diameter		diameter
129	Uniform and semi-	2.6 µm	Uniform and semi-	2.8 μm	30	151	Uniform and transparent		Uniform and transparent	1.2 μm
130	transparent Uniform and transparent	1.7 µm	transparent Uniform and transparent	1.9 μm		152	Uniform and semi- transparent	l 2.3 μm	Uniform and semi- transparent	2.6 μm
131	Uniform and transparent	1.2 μm	Uniform and transparent	1.4 μm	2.5	153	Uniform and transparent	l 0.9 μm	Uniform and transparent	1.1 μm
132	Uniform blue white	4.3 μm	Uniform blue white	4.6 µm	35	154	Uniform and semi-	2.6 μm	Uniform and semi-	2.9 µm
133	Uniform and semi- transparent	3.6 µm	Uniform and semi- transparent	3.9 µm		155	transparent Uniform and transparent	l 1.4 μm	transparent Uniform and transparent	1.5 μm
134	Uniform and semi-	3.3 µm	Uniform and semi-	3.5 µm	40	156	Uniform and transparent	l 1.3 μm	Uniform and transparent	1.4 µm
135	transparent Uniform and semi-	3.1 µm	transparent Uniform and semi-	3.3 µm		157	Uniform and transparent		Uniform and transparent	2.8 μm
136	transparent Uniform and	3.7 µm	transparent Uniform and	3.8 µm		158 159	Uniform and transparent Uniform and	·	Uniform and transparent Uniform and	1.3 μm 2.4 μm
	semi- transparent	·	semi- transparent	·	45	133	semi- transparent	2.2 μ	semi- transparent	2.1 pari
137	Uniform and transparent		Uniform and transparent	1.6 μm		160	Uniform blu white	·	Uniform blue white	4.5 μm
138 139	Uniform and transparent Uniform and	1.2 μm 2.5 μm	Uniform and transparent Uniform and	1.3 μm 2.7 μm	50	161	Uniform and semi- transparent	l 3.5 μm	Uniform and semi- transparent	3.8 µm
137	semi- transparent	·	semi- transparent	2./ pmi	50	162	Uniform and semi-	3.3 μm	Uniform and semi-	3.5 µm
140	Uniform and transparent	0.9 μm	Uniform and transparent	1.0 μm		163	transparent Uniform and	3.0 μm	transparent Uniform and	3.2 µm
141	Uniform and semi- transparent	2.2 µm	Uniform and semi- transparent	2.4 μm	55	164	semi- transparent	3.5	semi- transparent	2.0
142	Uniform and transparent	1.7 µm	Uniform and transparent	1.9 μm		164	Uniform and semi- transparent	l 3.5 µm	Uniform and semi- transparent	3.9 µm
143	Uniform and semi-	2.3 µm	Uniform and semi-	2.6 μm		165	Uniform and semi-	3.6 µm	Uniform and semi-	3.8 µm
144	transparent Uniform and transparent	1.9 µm	transparent Uniform and transparent	2.1 μm	60	166	transparent Uniform and transparent	l 1.7 μm	transparent Uniform and transparent	1.9 µm
145	Uniform and semi-	2.6 µm	Uniform and semi-	2.9 μm		167	Uniform and transparent	l 1.2 μm	Uniform and transparent	1.4 μm
146	transparent Uniform and	2.8 µm	transparent Uniform and	3.0 µm		168	Uniform and transparent		Uniform and transparent	1.7 µm
	semi- transparent		semi- transparent		65	169	Uniform and transparent	l 1.3 μm	Uniform and transparent	1.6 µm

61 TABLE 15-continued

62 TABLE 15-continued

		Evaluation of	solution stability		-			Evaluation of	solution stability	
				21	_					11
	Immed after pre		Leaving for 2		_ 5		Immed after pre		Leaving for 2	
Example	Visual observation	Average particle diameter	Visual observation	Average particle diameter		Example	Visual observation	Average particle diameter	Visual observation	Average particle diamete:
170	Uniform and	2.6 µm	Uniform and	2.9 µm	- 10	200	Uniform and	0.9 µm	Uniform and	1.0 µm
170	semi-	2.0 µm	semi-	2.9 μπ	10		transparent Uniform and		transparent	
171	transparent Uniform and	0.9 μm	transparent Uniform and	1.1 μm		201	semi-	2.7 µm	Uniform and semi-	2.9 μm
172	transparent Uniform and	1.8 µm	transparent Uniform and	1.9 µm		202	transparent Uniform and	1.8 µm	transparent Uniform and	1.9 µm
173	transparent Uniform and	1.6 µm	transparent Uniform and	1.8 µm	15	203	transparent Uniform and	1.7 µm	transparent Uniform and	1.8 µm
174	transparent Uniform and	1.4 μm	transparent Uniform and	1.6 μm		204	transparent Uniform and	1.2 μm	transparent Uniform and	1.4 μm
175	transparent Uniform and	1.5 µm	transparent Uniform and	1.6 µm		205	transparent Uniform and	1.5 µm	transparent Uniform and	1.6 µm
176	transparent Uniform and	1.0 µm	transparent Uniform and	1.2 μm	20	206	transparent Uniform and	1.6 μm	transparent Uniform and	1.7 μm
	transparent		transparent	·			transparent	,	transparent	
177	Uniform and semi-	2.4 μm	Uniform and semi-	2.7 μm		207	Uniform and transparent	1.2 μm	Uniform and transparent	1.3 μm
178	transparent Uniform and	0.8 μm	transparent Uniform and	0.9 μm		208	Uniform and transparent	1.0 μm	Uniform and transparent	1.1 μm
179	transparent Uniform and	1.4 μm	transparent Uniform and	1.6 µm	25	209	Uniform and transparent	1.1 µm	Uniform and transparent	1.2 μm
180	transparent Uniform and	1.8 µm	transparent Uniform and	2.0 μm		210	Uniform and transparent	1.3 µm	Uniform and transparent	1.5 μm
181	transparent Uniform and	1.1 μm	transparent Uniform and	1.3 μm		211	Uniform and transparent	0.9 μm	Uniform and transparent	1.0 μm
182	transparent Uniform and	2.7 μm	transparent Uniform and	3.0 μm	30	212	Uniform and transparent	1.9 µm	Uniform and transparent	2.1 μm
102	semi-	2.7 µm	semi-	3.0 µm		213	Uniform and semi-	2.5 μm	Uniform and semi-	2.7 μm
183	transparent Uniform and	1.9 µm	transparent Uniform and	2.0 μm		24.4	transparent	4.5	transparent	4.0
184	transparent Uniform and	1.8 μm	transparent Uniform and	1.9 µm	35	214	Uniform and transparent	1.7 μm	Uniform and transparent	1.9 µm
185	transparent Uniform and	2.4 µm	transparent Uniform and	2.7 µm		215	Uniform and semi-	2.4 µm	Uniform and semi-	2.6 µm
186	transparent Uniform and	1.3 µm	transparent Uniform and	1.4 μm		216	transparent Uniform and	1.0 µm	transparent Uniform and	1.1 μm
187	transparent Uniform and	1.6 µm	transparent Uniform and	1.7 µm		217	transparent Uniform and	1.2 µm	transparent Uniform and	1.4 μm
188	transparent Uniform and	2.5 µm	transparent Uniform and	2.7 μm	40		transparent		transparent	
100	semi- transparent	213 part	semi- transparent	217 pari		218	Uniform and transparent	1.4 µm	Uniform and transparent	1.6 µm
189	Uniform blue	4.1 μm	Uniform blue	4.4 µm		219	Uniform blue white	4.3 μm	Uniform blue white	4.8 µm
190	white Uniform and	3.8 µm	white Uniform and	3.9 µm	45	220	Uniform and semi-	3.8 µm	Uniform and semi-	4.0 μm
	semi- transparent		semi- transparent				transparent		transparent	
191	Uniform and semi-	3.6 µm	Uniform and semi-	3.8 µm		221	Uniform and semi-	3.4 µm	Uniform and semi-	3.6 µm
192	transparent Uniform and	3.4 µm	transparent Uniform and	3.5 µm	50	222	transparent Uniform and	3.2 µm	transparent Uniform and	3.5 µm
	semi- transparent	01. par	semi- transparent	0.10 pass	30		semi- transparent		semi- transparent	
193	Uniform and	3.2 µm	Uniform and	3.4 µm		223	Uniform and	3.3 µm	Uniform and	3.5 μm
	semi- transparent		semi- transparent				semi- transparent		semi- transparent	
194	Uniform and semi-	3.7 µm	Uniform and semi-	3.9 µm	55	224	Uniform and semi-	3.4 µm	Uniform and semi-	3.6 µm
195	transparent Uniform and	3.5 µm	transparent Uniform and	3.8 µm		225	transparent Uniform and	1.5 μm	transparent Uniform and	1.7 μm
170	semi-	<i>5.5</i> μm	semi-	5.6 µm			transparent	,	transparent	
196	transparent Uniform and	1.2 μm	transparent Uniform and	1.4 μm	60	226	Uniform and transparent	1.3 µm	Uniform and transparent	1.4 μm
197	transparent Uniform and	1.5 µm	transparent Uniform and	1.6 µm		227	Uniform and transparent	1.8 µm	Uniform and transparent	2.0 μm
198	transparent Uniform and	1.3 μm	transparent Uniform and	1.4 μm		228	Uniform and transparent	1.2 μm	Uniform and transparent	1.3 µm
	transparent	,	transparent	·	65	229	Uniform and	2.8 µm	Uniform and	3.0 μm
199	Uniform and transparent	1.7 µm	Uniform and transparent	1.8 µm	65		semi- transparent		semi- transparent	

63 TABLE 15-continued

64 TABLE 15-continued

		Evaluation of	solution stability		_			Evaluation of	solution stability	
	Immed after prep		Leaving for 2		_ 5		Immed after pre		Leaving for 2	
Example	Visual observation	Average particle diameter	Visual observation	Average particle diameter		Example	Visual observation	Average particle diameter	Visual observation	Average particle diameter
230	Uniform and semi-	2.5 µm	Uniform and semi-	2.7 µm	10	260	Uniform and transparent	1.7 µm	Uniform and transparent	1.8 µm
231	transparent Uniform and	1.1 μm	transparent Uniform and	1.2 μm		261	Uniform and transparent	1.6 µm	Uniform and transparent	1.7 µm
232	transparent Uniform and transparent	1.4 μm	transparent Uniform and transparent	1.6 µm		262 263	Uniform and transparent Uniform and	1.4 µm 1.2 µm	Uniform and transparent Uniform and	1.5 μm 1.4 μm
233	Uniform and semi-	2.6 µm	Uniform and semi-	2.8 µm	15	264	transparent Uniform and	2.1 µm	transparent Uniform and	2.4 µm
234	transparent Uniform and	0.9 μm	transparent Uniform and	1.0 μm			semi- transparent	·	semi- transparent	·
235	transparent Uniform and	1.4 μm	transparent Uniform and	1.5 µm	20	265	Uniform and semi-	2.7 µm	Uniform and semi-	3.0 µm
236	transparent Uniform and semi-	2.7 μm	transparent Uniform and semi-	2.9 μm		266	transparent Uniform and transparent	1.8 µm	transparent Uniform and transparent	2.0 μm
237	transparent Uniform and	1.2 μm	transparent Uniform and	1.4 μm		267	Uniform and transparent	1.9 μm	Uniform and transparent	2.0 μm
238	transparent Uniform and	2.3 μm	transparent Uniform and	2.5 μm	25	268	Uniform and transparent	1.1 μm	Uniform and transparent	1.2 μm
239	semi- transparent Uniform and	1.1 μm	semi- transparent Uniform and	1.2 μm		269	Uniform and semi- transparent	2.3 µm	Uniform and semi- transparent	2.4 μm
240	transparent Uniform and	2.6 μm	transparent Uniform and	2.8 μm		270	Uniform and semi-	2.7 µm	Uniform and semi-	2.9 µm
241	semi- transparent	1.5	semi- transparent	1.6	30	271	transparent Uniform blue	4.5 μm	transparent Uniform blue	4.8 μm
241 242	Uniform and transparent Uniform and	1.5 µm 1.7 µm	Uniform and transparent Uniform and	1.6 μm 1.8 μm		272	white Uniform and transparent	1.1 μm	white Uniform and transparent	1.2 μm
243	transparent Uniform and	2.4 μm	transparent Uniform and	2.6 µm	35	273	Uniform and transparent	1.5 µm	Uniform and transparent	1.7 µm
	semi- transparent	·	semi- transparent	·	33	274	Uniform and transparent	1.3 µm	Uniform and transparent	1.4 μm
244	Uniform and transparent	1.1 μm	Uniform and transparent	1.2 μm		275	Uniform and transparent	1.5 μm	Uniform and transparent	1.6 μm
245	Uniform and semi- transparent	2.6 µm	Uniform and semi- transparent	2.9 µm	40	276 277	Uniform and transparent Uniform and	0.9 µm 2.4 µm	Uniform and transparent Uniform and	1.0 µm 2.7 µm
246	Uniform and transparent	1.3 µm	Uniform and transparent	1.4 µm		277	semi- transparent	2.+ μπ	semi- transparent	2.7 µm
247	Uniform and transparent	1.3 μm	Uniform and transparent	1.4 μm		278	Uniform and transparent	1.7 µm	Uniform and transparent	1.8 µm
248	Uniform blue white	4.4 μm	Uniform blue white	4.8 μm	45	279	Uniform and transparent	1.8 µm	Uniform and transparent	2.0 μm
249 250	Uniform and transparent Uniform and	1.7 μm 1.5 μm	Uniform and transparent Uniform and	1.9 μm 1.6 μm		280 281	Uniform and transparent Uniform and	1.5 μm 2.7 μm	Uniform and transparent Uniform and	1.7 µm 2.9 µm
251	transparent Uniform and	1.1 μm	transparent Uniform and	1.2 µm		201	semi- transparent	2.7 µm	semi- transparent	2.5 μπ
252	transparent Uniform and	1.8 μm	transparent Uniform and	2.μm	50	282	Uniform and transparent	1.4 μm	Uniform and transparent	1.5 μm
253	transparent Uniform and	2.5 μm	transparent Uniform and	2.8 μm		283	Uniform and transparent	1.4 μm	Uniform and transparent	1.5 μm
254	semi- transparent	1.2	semi- transparent	1.4		284 285	Uniform and transparent Uniform and	1.2 µm 2.2 µm	Uniform and transparent Uniform and	1.4 μm 2.5 μm
254 255	Uniform and transparent Uniform and	1.3 μm 1.1 μm	Uniform and transparent Uniform and	1.4 μm 1.2 μm	55	200	semi- transparent	2.2 part	semi- transparent	2.0 pm
256	transparent Uniform and	1.1 μm 1.0 μm	transparent Uniform and	1.2 μm 1.1 μm		286	Uniform and transparent	1.5 μm	Uniform and transparent	1.6 μm
257	transparent Uniform and	2.7 μm	transparent Uniform and	2.9 μm	60	287 288	Uniform and transparent Uniform and	1.7 µm 2.3 µm	Uniform and transparent Uniform and	1.4 μm 2.5 μm
	semi- transparent	·	semi- transparent	·		200	semi- transparent	2.5 µm	semi- transparent	2.5 µm
258	Uniform and semi-	2.3 μm	Uniform and semi-	2.6 µm		289	Uniform and transparent	1.3 μm	Uniform and transparent	1.5 μm
	transparent Uniform and	1.5 µm	transparent Uniform and	1.6 µm	65	290	Uniform and semi-	2.5 µm	Uniform and semi-	2.8 µm

TABLE 15-continued

		Evaluation of	solution stability	
	Immed after prep		Leaving for 2	
Example	Visual observation	Average particle diameter	Visual observation	Average particle diameter
291	Uniform and semi-	2.8 µm	Uniform and semi-	3.0 µm
292	transparent Uniform and semi-	2.4 μm	transparent Uniform and semi-	2.6 μm
293	transparent Uniform and semi-	2.1 μm	transparent Uniform and semi-	2.3 μm
294	transparent Uniform and semi-	2.9 μm	transparent Uniform and semi-	3.0 μm
295	transparent Uniform and transparent	1.2 μm	transparent Uniform and transparent	1.5 μm
296	Uniform blue white	4.2 µm	Uniform blue white	4.6 μm
297	Uniform and	0.8 μm	Uniform and	1.0 μm
298	transparent Uniform and transparent	0.9 μm	transparent Uniform and transparent	1.1 μm
299	Uniform and	0.9 μm	Uniform and	1.1 μm
300	transparent Uniform and	0. 8 μm	transparent Uniform and	1.0 μm
701	transparent Uniform and transparent	0.9 μm	transparent Uniform and transparent	1.0 μm

TABLE 16

	E	valuation of	solution stability	
	Immediat prepar		Leaving for	
Comparative Example	Visual observation	Average particle diameter	Visual observation	Average particle diameter
Comparative Example 1	Sedimented, coalesced	17.7 μm	Sedimented, coalesced	76.2 µm
Comparative Example 2	Sedimented, coalesced	16.9 µm	Sedimented, coalesced	78.3 µm
Comparative Example 3	Sedimented, coalesced	19.2 µm	Sedimented, coalesced	76.7 µm
Comparative Example 4	Sedimented, coalesced	16.8 µm	Sedimented, coalesced	82.4 µm
Comparative Example 5	Sedimented, coalesced	15.7 μm	Sedimented, coalesced	75.7 µm
Comparative Example 6	Sedimented, coalesced	16.5 μm	Sedimented, coalesced	76.4 µm

In Tables 7 to 13, each of the contents of the fluorine-atom-containing polyacrylate, the fluorine-atom-containing polymethacrylate, the polycarbonate having a siloxane bond, the polystyrene having a siloxane bond, the compound represented by the formula (5), the compound represented by the formula (6), and the compound represented by the formula (7) is a content thereof based on the charge transporting substance and binder resin (% by mass).

By comparison of Examples with Comparative Examples, in the production method in which the solution containing the charge transporting substance and at least one compound 65 selected from the group consisting of the fluorine-atom-containing polyacrylate, the fluorine-atom-containing

polymethacrylate, the polycarbonate having a siloxane bond, the polyester having a siloxane bond, the polystyrene having a siloxane bond, the silicone oil, the polyolefin, the aliphatic acid, the aliphatic acid amide, and the aliphatic acid ester is prepared, and the emulsion is prepared using the solution and water, the state of the emulsion is stably kept during preservation for a long time, and the same state of that of the emulsion immediately after preparation is kept. In the conventional emulsion described in Japanese Patent Application Laid-Open No. 2011-128213, however, by addition of the surfactant, the oil droplets containing the charge transporting substance and the binder resin are relatively stable immediately after the emulsion is prepared, but the oil droplets may coalesce after long-term preservation, leading to aggregation. A method for increasing the content of the surfactant to suppress coalescence is thought, but usually, the surfactant easily results in reduction in the electrophotographic properties. Accordingly, the method is not con-20 sidered desirable.

In the method according to the present invention in which the solution containing the charge transporting substance and the compound that reduces the surface energy is prepared, and the emulsion is prepared, the compound that reduces the surface energy exists on the surfaces of the oil droplets. For this reason, the surface energy can be reduced, and occurrence of aggregation of the oil droplets can be significantly suppressed compared to the case where the compound that reduces the surface energy is not used. This method provides long-term solution stability of the emulsion, and the emulsion is useful as the coating solution for the electrophotographic photosensitive member.

An aluminum cylinder having a diameter of 30 mm and a length of 260.5 mm was used as the support (electrically conductive support). Next, 10 parts of SnO₂ coated barium sulfate (conductive particle), 2 parts of titanium oxide (pigment for adjusting resistance), 6 parts of a phenol resin, and 0.001 parts of a silicone oil (leveling agent) were dissolved using a mixed solvent of 4 parts of methanol and 16 parts of methoxypropanol to prepare a coating solution for an electrically conductive layer. The coating solution for an electrically conductive layer was applied onto the aluminum cylinder by dip coating. The obtained coat was cured (thermally cured) at 140° C. for 30 minutes to form an electrically conductive layer having a film thickness of 15 μm.

Next, 3 parts of N-methoxymethylated nylon and 3 parts of a copolymerized nylon were dissolved in a mixed solvent of 65 parts of methanol and 30 parts of n-butanol to prepare a coating solution for an undercoat layer. The coating solution for an undercoat layer was applied onto the electrically conductive layer by dip coating. The obtained coat was dried at 100° C. for 10 minutes to form an undercoat layer having a film thickness of 0.7 μm.

Next, 10 parts of a crystalline hydroxy gallium phthalocyanine (charge generating substance) having strong peaks at Bragg angles $(20\pm0.2^{\circ})$ of 7.5° , 9.9° , 16.3° , 18.6° , 25.1° , and 28.3° in CuK α properties X ray diffraction was prepared. 250 parts of cyclohexanone and 5 parts of a polyvinyl butyral (trade name: S-LEC BX-1, made by Sekisui Chemical Co., Ltd.) were mixed with the hydroxy gallium phthalocyanine, and dispersed for 1 hour under an atmosphere of $23\pm3^{\circ}$ C. using a sand mill apparatus having glass beads whose diameter was 1 mm. After dispersion, 250 parts of ethyl acetate was added to prepare a coating solution for a charge generating layer. The coating solution for a charge generating layer was applied onto the undercoat layer by dip

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coating. The obtained coat was dried at 100° C. for 10 minutes to form a charge generating layer having a film thickness of $0.26~\mu m$.

Next, as the coating solution for a charge transporting layer, the emulsion prepared in Example 1 was applied onto the charge generating layer by dip coating to form a coat of the emulsion. The obtained coat was heated at 130° C. for 1 hour to form a charge transporting layer having a film thickness of $20~\mu m$. Thus, an electrophotographic photosensitive member was produced. The used emulsion and the heating condition for the coat formed by applying the emulsion are shown in Table 17. The emulsion used for dip coating was left as it was for 2 weeks (under an environment of the temperature of 23° C. and humidity of 50% RH), and stirred at 1,000 turns/min for 3 minutes by a homogenizer.

Next, evaluations will be described.

<Evaluation of Uniformity of Coat (Coat Uniformity)>

A place 130 mm from the upper end of the surface of the electrophotographic photosensitive member was measured using a surface roughness measuring apparatus (SURF-CORDER SE-3400, made by Kosaka Laboratory Ltd.), and evaluation was made according to evaluation of the tenpoint height of irregularities (Rzjis) according to JIS B 0601:2001 (evaluation length of 10 mm). The results are shown in Table 17.

<Evaluation of Image>

In a laser beam printer LBP-2510 made by Canon Inc., the charge potential (dark potential) of the electrophotographic photosensitive member and the exposure amount (image exposure amount) of a laser light source at 780 nm were modified such that the light amount on the surface of the electrophotographic photosensitive member was $0.3~\mu\text{J/cm}^2$. The thus-modified laser beam printer LBP-2510 was used. Evaluation was made under an environment of the temperature of 23° C. and relative humidity of 15% RH. In evaluation of an image, an A4 size normal paper was used, and a halftone image of a single color was output. The output image was visually evaluated on the criterion below. The results are shown in Table 17.

Rank A: a totally uniform image is found

Rank B: very slight unevenness is found in an image

Rank C: unevenness is found in an image

Rank D: remarkable unevenness is found in an image

Examples 302 to 600

An electrophotographic photosensitive member was produced by the same method as that in Example 301 except that the emulsion used in formation of the charge transporting layer was changed to the emulsion shown in Tables 17 and 18. The electrophotographic photosensitive member 50 was evaluated by the same method as that in Example 301. The results are shown in Tables 17 and 18.

Example 801

An electrophotographic photosensitive member was produced by the same method as that in Example 301 except that the emulsion used in formation of the charge transporting layer was changed to the emulsion described in Example 701. The electrophotographic photosensitive member was evaluated by the same method as that in Example 301. The results are shown in Table 18.

Comparative Examples 7 to 12

An electrophotographic photosensitive member was produced by the same method as that in Example 301 except

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that the emulsion used in formation of the charge transporting layer was changed to the emulsion shown in Table 19. The electrophotographic photosensitive member was evaluated by the same method as that in Example 301. The results are shown in Table 19. Gentle depressions and projections were formed on the obtained electrophotographic photosensitive member, and unevenness of the image corresponding to the depressions and projections was detected as the image.

Comparative Examples 13 and 14

An electrophotographic photosensitive member was produced by the same method as that in Example 301 except that the prepared emulsion was not left for 2 weeks in Example 301, and was immediately applied by dip coating, the emulsion was used in formation shown in Table 19, and the heating condition for the coat formed by applying the emulsion was changed as shown in Table 19. The electrophotographic photosensitive member was evaluated by the same method as that in Example 301. The results are shown in Table 19. Gentle depressions and projections were formed on the obtained electrophotographic photosensitive member, and unevenness of the image corresponding to the depressions and projections was detected as the image.

TABLE 17

•			Heating	condition	Evaluation of	
	Exam- ple	Emulsion	Temper- ature	Time	uniformity of coat	Evaluation of image
	301	Example 1	130° C.	60 Minutes	0.49	A
	302	Example 2	130° C.	60 Minutes	0.57	A
	303	Example 3	130° C.	60 Minutes	0.60	Α
	304	Example 4	130° C.	60 Minutes	0.45	A
	305	Example 5	130° C.	60 Minutes	0.55	A
	306	Example 6	130° C.	60 Minutes	0.50	A
	307	Example 7	130° C.	60 Minutes	0.47	A
	308	Example 8	130° C.	60 Minutes	0.49	A
	309	Example 9	130° C.	60 Minutes	0.49	A
1	310	Example 10	130° C.	60 Minutes	0.54	A
	311	Example 11	130° C.	60 Minutes	0.50	A
	312	Example 12	130° C.	60 Minutes	0.46	A
	313	Example 13	130° C.	60 Minutes	0.48	A
	314	Example 14	130° C.	60 Minutes	0.58	A
	315	Example 15	130° C. 130° C.	60 Minutes 60 Minutes	0.59	A A
	316 317	Example 16	130° C.	60 Minutes	0.55 0.56	A A
	318	Example 17 Example 18	130° C.	60 Minutes	0.50	A A
	319	Example 19	130° C.	60 Minutes	0.32	A A
	320	Example 19	130° C.	60 Minutes	0.49	A
	321	Example 20	130° C.	60 Minutes	0.50	A
	322	Example 22	130° C.	60 Minutes	0.51	A
	323	Example 23	130° C.	60 Minutes	0.57	A
	324	Example 24	130° C.	60 Minutes	0.66	В
	325	Example 25	130° C.	60 Minutes	0.68	A
	326	Example 26	130° C.	60 Minutes	0.68	A
	327	Example 27	130° C.	60 Minutes	0.66	В
	328	Example 28	130° C.	60 Minutes	0.62	A
	329	Example 29	130° C.	60 Minutes	0.68	A
	330	Example 30	130° C.	60 Minutes	0.68	Ā
	331	Example 31	130° C.	60 Minutes	0.68	A
	332	Example 32	130° C.	60 Minutes	0.67	A
	333	Example 33	130° C.	60 Minutes	0.69	A
	334	Example 34	130° C.	60 Minutes	0.69	A
	335	Example 35	130° C.	60 Minutes	0.51	A
1	336	Example 36	130° C.	60 Minutes	0.59	A
	337	Example 37	130° C.	60 Minutes	0.50	A
	338	Example 38	130° C.	60 Minutes	0.57	A
	339	Example 39	130° C.	60 Minutes	0.46	A
	340	Example 40	130° C.	60 Minutes	0.58	\mathbf{A}
	341	Example 41	130° C.	60 Minutes	0.50	A
	342	Example 42	130° C.	60 Minutes	0.60	A
	343	Example 43	130° C.	60 Minutes	0.48	A

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TABLE 17-continued

70 TABLE 17-continued

			/-continued	•					IADLE I	/-continued	•	
		Heating	condition	Evaluation of					Heating	g condition	Evaluation of	
Exam- ple	Emulsion	Temper- ature	Time	uniformity of coat	Evaluation of image	5	Exam- ple	Emulsion	Temper- ature	Time	uniformity of coat	Evaluation of image
344	Example 44	130° C.	60 Minutes	0.49	A		418	Example 118	130° C.	60 Minutes	0.60	A
345	Example 45	130° C.	60 Minutes	0.55	A		419	Example 119	130° C.	60 Minutes	0.48	A
346 347	Example 46 Example 47	130° C. 130° C.	60 Minutes 60 Minutes	0.47 0.46	A	10	420 421	Example 120	130° C. 130° C.	60 Minutes 60 Minutes	0.55	A
347	Example 47	130° C.	60 Minutes	0.46	A A	10	421	Example 121 Example 122	130° C.	60 Minutes	0.47 0.48	A A
349	Example 49	130° C.	60 Minutes	0.56	A		423	Example 123	130° C.	60 Minutes	0.59	A
350	Example 50	130° C.	60 Minutes	0.55	A		424	Example 124	130° C.	60 Minutes	0.56	A
351	Example 51	130° C.	60 Minutes	0.52	A		425	Example 125	130° C.	60 Minutes	0.57	\mathbf{A}
352 353	Example 52 Example 53	130° C. 130° C.	60 Minutes 60 Minutes	0.55 0.49	A A		426	Example 126	130° C.	60 Minutes	0.49	A
354	Example 54	130° C.	60 Minutes	0.54	A	15	427	Example 127	130° C.	60 Minutes	0.48	A
355	Example 55	130° C.	60 Minutes	0.49	A		428 429	Example 128 Example 129	130° C. 130° C.	60 Minutes 60 Minutes	0.47 0.52	A A
356	Example 56	130° C.	60 Minutes	0.48	A		430	Example 129	130° C.	60 Minutes	0.54	A
357	Example 57	130° C.	60 Minutes	0.47	A		431	Example 131	130° C.	60 Minutes	0.68	В
358 359	Example 58 Example 59	130° C. 130° C.	60 Minutes 60 Minutes	0.51 0.56	A A		432	Example 132	130° C.	60 Minutes	0.61	A
360	Example 60	130° C.	60 Minutes	0.52	A	20	433	Example 133	130° C.	60 Minutes	0.63	A
361	Example 61	130° C.	60 Minutes	0.59	A		434	Example 134	130° C.	60 Minutes	0.66	В
362	Example 62	130° C.	60 Minutes	0.58	A		435	Example 135	130° C.	60 Minutes	0.68	A
363	Example 63	130° C.	60 Minutes	0.58	A		436	Example 136	130° C.	60 Minutes	0.58	A
364 365	Example 64 Example 65	130° C. 130° C.	60 Minutes 60 Minutes	0.54 0.57	A A		437 438	Example 137 Example 138	130° C. 130° C.	60 Minutes 60 Minutes	0.51 0.49	A A
366	Example 66	130° C.	60 Minutes	0.60	A	25	439	Example 139	130° C.	60 Minutes	0.58	A
367	Example 67	130° C.	60 Minutes	0.48	A		440	Example 140	130° C.	60 Minutes	0.60	A
368	Example 68	130° C.	60 Minutes	0.46	A		441	Example 141	130° C.	60 Minutes	0.57	A
369	Example 69	130° C.	60 Minutes	0.54	A		442	Example 142	130° C.	60 Minutes	0.59	A
370	Example 70	130° C.	60 Minutes 60 Minutes	0.54	A		443	Example 143	130° C.	60 Minutes	0.59	A
371 372	Example 71 Example 72	130° C. 130° C.	60 Minutes	0.52 0.47	A A	30	444	Example 144	130° C.	60 Minutes	0.47	A
373	Example 73	130° C.	60 Minutes	0.54	A	30	445	Example 145	130° C.	60 Minutes	0.57	A
374	Example 74	130° C.	60 Minutes	0.46	A		446	Example 146	130° C.	60 Minutes	0.51	A
375	Example 75	130° C.	60 Minutes	0.52	A		447 448	Example 147 Example 148	130° C. 130° C.	60 Minutes 60 Minutes	0.50 0.46	A A
376	Example 76	130° C.	60 Minutes	0.54	A		449	Example 149	130° C.	60 Minutes	0.52	A
377 378	Example 77 Example 78	130° C. 130° C.	60 Minutes 60 Minutes	0.50 0.58	A A		450	Example 150	130° C.	60 Minutes	0.52	A
379	Example 79	130° C.	60 Minutes	0.66	В	35						
380	Example 80	130° C.	60 Minutes	0.48	A							
381	Example 81	130° C.	60 Minutes	0.57	A							
382	Example 82											
		130° C.	60 Minutes	0.57	A				TAB	LE 18		
383 384	Example 83 Example 84	130° C.	60 Minutes	0.59	A				TAB	BLE 18	T. I. di	
384 385	Example 83 Example 84 Example 85					40					Evaluation	
384	Example 84	130° C. 130° C.	60 Minutes 60 Minutes	0.59 0.52	A A	40				LE 18	Evaluation of	
384 385 386 387	Example 84 Example 85 Example 86 Example 87	130° C. 130° C. 130° C. 130° C. 130° C.	60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes	0.59 0.52 0.46 0.51 0.58	A A A A	40	Exam-				of of	Evaluation
384 385 386 387 388	Example 84 Example 85 Example 86 Example 87 Example 88	130° C. 130° C. 130° C. 130° C. 130° C. 130° C.	60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59	A A A A A	40	Example	Emulsion	Heating		of of	Evaluation of image
384 385 386 387 388 389	Example 84 Example 85 Example 86 Example 87 Example 88 Example 89	130° C. 130° C. 130° C. 130° C. 130° C. 130° C. 130° C.	60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59	A A A A A A	40	ple		Heating Temper- ature	g condition Time	of uniformity of coat	of image
384 385 386 387 388	Example 84 Example 85 Example 86 Example 87 Example 88 Example 89 Example 90	130° C. 130° C. 130° C. 130° C. 130° C. 130° C.	60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59	A A A A A	40		Example	Heating Temper-	g condition	of uniformity	
384 385 386 387 388 389 390	Example 84 Example 85 Example 86 Example 87 Example 88 Example 89	130° C. 130° C. 130° C. 130° C. 130° C. 130° C. 130° C. 130° C.	60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54	A A A A A A A		9le 451	Example 151	Heating Temperature 130° C.	Time 60 Minutes	of uniformity of coat	of image
384 385 386 387 388 389 390 391 392 393	Example 84 Example 85 Example 86 Example 87 Example 88 Example 89 Example 90 Example 91 Example 92 Example 93	130° C. 130° C. 130° C. 130° C. 130° C. 130° C. 130° C. 130° C. 130° C. 130° C.	60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54 0.48 0.60 0.62	A A A A A A A A A		451 452	Example 151 Example 152	Heating Temperature 130° C. 130° C.	Time 60 Minutes 60 Minutes	of uniformity of coat 0.57 0.53	of image
384 385 386 387 388 389 390 391 392 393 394	Example 84 Example 85 Example 86 Example 87 Example 88 Example 89 Example 90 Example 91 Example 92 Example 93 Example 94	130° C. 130° C.	60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54 0.48 0.60 0.62 0.66	A A A A A A A A A B		9le 451	Example 151 Example 152 Example	Heating Temperature 130° C.	Time 60 Minutes	of uniformity of coat	of image
384 385 386 387 388 389 390 391 392 393 394 395	Example 84 Example 85 Example 86 Example 87 Example 88 Example 89 Example 90 Example 91 Example 92 Example 93 Example 94 Example 95	130° C. 130° C.	60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54 0.48 0.60 0.62 0.66 0.63	A A A A A A A A A A A A A A A A A A A		451 452 453	Example 151 Example 152 Example 153	Heating Temper- ature 130° C. 130° C. 130° C.	Time 60 Minutes 60 Minutes 60 Minutes	of uniformity of coat 0.57 0.53 0.53	A A A
384 385 386 387 388 389 390 391 392 393 394 395 396	Example 84 Example 85 Example 86 Example 87 Example 88 Example 89 Example 90 Example 91 Example 92 Example 93 Example 94 Example 95 Example 96	130° C. 130° C.	60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54 0.48 0.60 0.62 0.63 0.69	A A A A A A A A A A A A A A A A A A A	45	451 452	Example 151 Example 152 Example 153 Example	Heating Temperature 130° C. 130° C.	Time 60 Minutes 60 Minutes	of uniformity of coat 0.57 0.53	of image A A
384 385 386 387 388 389 390 391 392 393 394 395	Example 84 Example 85 Example 86 Example 87 Example 88 Example 89 Example 90 Example 91 Example 92 Example 93 Example 94 Example 95	130° C. 130° C.	60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54 0.48 0.60 0.62 0.66 0.63	A A A A A A A A A A A A A A A A A A A		451 452 453	Example 151 Example 152 Example 153	Temper- ature 130° C. 130° C. 130° C. 130° C.	Time 60 Minutes 60 Minutes 60 Minutes 60 Minutes	of uniformity of coat 0.57 0.53 0.53 0.46	A A A A
384 385 386 387 388 389 390 391 392 393 394 395 396 397 398 399	Example 84 Example 85 Example 86 Example 87 Example 88 Example 90 Example 90 Example 91 Example 92 Example 93 Example 94 Example 95 Example 96 Example 97 Example 97 Example 98 Example 99	130° C. 130° C.	60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54 0.60 0.62 0.66 0.63 0.69 0.51 0.55 0.55	A A A A A A A A B A A A A A A A A A A A	45	ple 451 452 453 454	Example 151 Example 152 Example 153 Example 154	Heating Temper- ature 130° C. 130° C. 130° C.	Time 60 Minutes 60 Minutes 60 Minutes	of uniformity of coat 0.57 0.53 0.53	A A A
384 385 386 387 388 390 391 392 393 394 395 396 397 398 399 400	Example 84 Example 85 Example 86 Example 87 Example 88 Example 90 Example 91 Example 92 Example 93 Example 94 Example 95 Example 96 Example 97 Example 97 Example 98 Example 99 Example 100	130° C.	60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54 0.60 0.62 0.66 0.63 0.69 0.51 0.55 0.58	A A A A A A A A A A A A A A A A A A A	45	ple 451 452 453 454	Example 151 Example 152 Example 153 Example 154 Example 155 Example	Temper- ature 130° C. 130° C. 130° C. 130° C.	Time 60 Minutes 60 Minutes 60 Minutes 60 Minutes	of uniformity of coat 0.57 0.53 0.53 0.46	A A A A
384 385 386 387 388 389 390 391 392 393 394 395 396 397 398 399 400 401	Example 84 Example 85 Example 86 Example 87 Example 89 Example 90 Example 91 Example 92 Example 93 Example 94 Example 95 Example 96 Example 97 Example 98 Example 99 Example 100 Example 100	130° C.	60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54 0.48 0.60 0.62 0.66 0.63 0.69 0.51 0.55 0.58	A A A A A A A A A A A A A A A A A A A	45	9le 451 452 453 454 455 456	Example 151 Example 152 Example 153 Example 154 Example 155 Example 155	Heating Temper- ature 130° C. 130° C. 130° C. 130° C. 130° C.	Time 60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes	of uniformity of coat 0.57 0.53 0.53 0.46 0.52 0.57	A A A A A A
384 385 386 387 388 389 390 391 392 393 394 395 396 397 398 399 400 401 402	Example 84 Example 85 Example 86 Example 87 Example 89 Example 90 Example 91 Example 92 Example 93 Example 94 Example 95 Example 96 Example 97 Example 98 Example 100 Example 101 Example 101	130° C.	60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54 0.48 0.60 0.62 0.66 0.63 0.69 0.51 0.55 0.58 0.51 0.55	A A A A A A A A A A A A A A A A A A A	45 50	ple 451 452 453 454 455	Example 151 Example 152 Example 153 Example 154 Example 155 Example 156 Example	Heating Temper- ature 130° C. 130° C. 130° C. 130° C.	Time 60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes 60 Minutes	of uniformity of coat 0.57 0.53 0.53 0.46 0.52	A A A A A
384 385 386 387 388 389 390 391 392 393 394 395 396 397 398 399 400 401	Example 84 Example 85 Example 86 Example 87 Example 89 Example 90 Example 91 Example 92 Example 93 Example 94 Example 95 Example 96 Example 97 Example 98 Example 99 Example 100 Example 100	130° C.	60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54 0.48 0.60 0.62 0.66 0.63 0.69 0.51 0.55 0.58	A A A A A A A A A A A A A A A A A A A	45	9le 451 452 453 454 455 456	Example 151 Example 152 Example 153 Example 154 Example 155 Example 155 Example 155 Example 156 Example 156	Heating Temper- ature 130° C. 130° C. 130° C. 130° C. 130° C.	Time 60 Minutes	of uniformity of coat 0.57 0.53 0.53 0.46 0.52 0.57 0.54	A A A A A A
384 385 386 387 388 389 390 391 392 393 394 395 396 397 398 399 400 401 402 403 404 405	Example 84 Example 85 Example 86 Example 88 Example 89 Example 90 Example 91 Example 92 Example 93 Example 95 Example 96 Example 97 Example 98 Example 100 Example 100 Example 101 Example 102 Example 103 Example 104 Example 104 Example 104 Example 105	130° C.	60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.48 0.60 0.62 0.66 0.63 0.69 0.51 0.55 0.58 0.59 0.51 0.55 0.58	A A A A A A A A A A A A A A A A A A A	45 50	ple 451 452 453 454 455 456 457 458	Example 151 Example 152 Example 153 Example 154 Example 155 Example 156 Example	Heating Temper- ature 130° C. 130° C. 130° C. 130° C. 130° C. 130° C.	Time 60 Minutes	of uniformity of coat 0.57 0.53 0.53 0.46 0.52 0.57	A A A A A A
384 385 386 387 388 389 390 391 392 393 394 395 396 397 398 399 400 401 402 403 404 405 406	Example 84 Example 85 Example 86 Example 87 Example 89 Example 90 Example 91 Example 92 Example 93 Example 94 Example 95 Example 96 Example 97 Example 98 Example 100 Example 101 Example 101 Example 102 Example 103 Example 105 Example 105 Example 105 Example 106	130° C.	60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54 0.48 0.60 0.62 0.66 0.63 0.69 0.51 0.55 0.58 0.51 0.50 0.58	A A A A A A A A A A A A A A A A A A A	45 50	ple 451 452 453 454 455 456 457	Example 151 Example 152 Example 153 Example 154 Example 155 Example 156 Example 157 Example 157 Example	Heating Temper- ature 130° C. 130° C. 130° C. 130° C. 130° C. 130° C.	Time 60 Minutes	of uniformity of coat 0.57 0.53 0.53 0.46 0.52 0.57 0.54	A A A A A A
384 385 386 387 388 389 390 391 392 393 394 395 396 397 398 399 400 401 402 403 404 405 406 407	Example 84 Example 85 Example 86 Example 87 Example 89 Example 90 Example 91 Example 93 Example 94 Example 94 Example 96 Example 97 Example 97 Example 100 Example 101 Example 101 Example 102 Example 104 Example 104 Example 104 Example 106 Example 106 Example 106 Example 107	130° C.	60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54 0.48 0.60 0.62 0.66 0.63 0.69 0.51 0.55 0.58 0.51 0.50 0.58 0.59	A A A A A A A A A A A A A A A A A A A	45 50	ple 451 452 453 454 455 456 457 458 459	Example 151 Example 152 Example 153 Example 154 Example 155 Example 156 Example 157 Example 157 Example 158 Example 158	Heating Temper- ature 130° C.	Time 60 Minutes	of uniformity of coat 0.57 0.53 0.53 0.46 0.52 0.57 0.54 0.56 0.46	A A A A A A A A
384 385 386 387 388 389 390 391 392 393 394 395 396 397 398 399 400 401 402 403 404 405 406 407 408	Example 84 Example 85 Example 86 Example 88 Example 89 Example 90 Example 91 Example 92 Example 93 Example 94 Example 95 Example 96 Example 97 Example 97 Example 100 Example 100 Example 101 Example 102 Example 102 Example 104 Example 105 Example 104 Example 105 Example 106 Example 107 Example 107 Example 107 Example 107	130° C.	60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54 0.48 0.60 0.62 0.66 0.63 0.69 0.51 0.55 0.58 0.51 0.50 0.58 0.51 0.59 0.51 0.55 0.53 0.59 0.51 0.55 0.54 0.60 0.62 0.63 0.69 0.51 0.55	A A A A A A A A A A A A A A A A A A A	45 50	ple 451 452 453 454 455 456 457 458	Example 151 Example 152 Example 153 Example 154 Example 155 Example 156 Example 157 Example 158 Example 158 Example 158 Example	Heating Temper- ature 130° C.	Time 60 Minutes	0.57 0.53 0.53 0.54 0.54 0.56	A A A A A A A
384 385 386 387 388 389 390 391 392 393 394 395 396 397 398 399 400 401 402 403 404 405 406 407	Example 84 Example 85 Example 86 Example 87 Example 89 Example 90 Example 91 Example 93 Example 94 Example 94 Example 96 Example 97 Example 97 Example 100 Example 101 Example 101 Example 102 Example 104 Example 104 Example 104 Example 106 Example 106 Example 106 Example 107	130° C.	60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54 0.48 0.60 0.62 0.66 0.63 0.69 0.51 0.55 0.58 0.51 0.50 0.58 0.59	A A A A A A A A A A A A A A A A A A A	45 50	ple 451 452 453 454 455 456 457 458 459 460	Example 151 Example 152 Example 153 Example 154 Example 155 Example 155 Example 156 Example 157 Example 157 Example 158 Example 158 Example 158 Example 159 Example 159 Example 160	Heating Temper- ature 130° C.	Time 60 Minutes	0.57 0.53 0.53 0.46 0.52 0.57 0.54 0.56 0.46 0.64	A A A A A A A B
384 385 386 387 388 389 390 391 392 393 394 395 396 397 398 399 400 401 402 403 404 405 406 407 408 409 410 411	Example 84 Example 85 Example 86 Example 87 Example 89 Example 90 Example 91 Example 93 Example 94 Example 95 Example 96 Example 97 Example 98 Example 100 Example 101 Example 102 Example 103 Example 104 Example 105 Example 106 Example 106 Example 107 Example 108 Example 108 Example 108 Example 108 Example 110 Example 111	130° C.	60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54 0.48 0.60 0.62 0.66 0.63 0.69 0.51 0.55 0.58 0.50 0.58 0.50 0.58 0.59 0.50 0.51 0.55	A A A A A A A A A A A A A A A A A A A	455055	ple 451 452 453 454 455 456 457 458 459	Example 151 Example 152 Example 153 Example 154 Example 155 Example 156 Example 157 Example 158 Example 158 Example 158 Example	Heating Temper- ature 130° C.	Time 60 Minutes	of uniformity of coat 0.57 0.53 0.53 0.46 0.52 0.57 0.54 0.56 0.46	A A A A A A A A
384 385 386 387 388 389 390 391 392 393 394 395 396 397 398 399 400 401 402 403 404 405 406 407 408 409 411 412	Example 84 Example 85 Example 86 Example 87 Example 89 Example 90 Example 91 Example 93 Example 94 Example 95 Example 96 Example 97 Example 97 Example 100 Example 101 Example 102 Example 104 Example 104 Example 104 Example 106 Example 106 Example 107 Example 107 Example 108 Example 109 Example 101 Example 111 Example 111	130° C.	60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54 0.48 0.60 0.62 0.66 0.63 0.69 0.51 0.55 0.58 0.51 0.50 0.58 0.56 0.46 0.51 0.55 0.58 0.51 0.55 0.58 0.51 0.55 0.58 0.51 0.55 0.58 0.51 0.55 0.58 0.51 0.55 0.56 0.56 0.57 0.50	A A A A A A A A A A A A A A A A A A A	455055	ple 451 452 453 454 455 456 457 458 459 460	Example 151 Example 152 Example 153 Example 154 Example 155 Example 156 Example 157 Example 158 Example 158 Example 160 Example 160 Example	Heating Temper- ature 130° C.	Time 60 Minutes	0.57 0.53 0.53 0.46 0.52 0.57 0.54 0.56 0.46 0.64	A A A A A A A B
384 385 386 387 388 389 390 391 392 393 394 395 396 397 398 399 400 401 402 403 404 405 406 407 408 409 410 411 412 413	Example 84 Example 85 Example 86 Example 88 Example 89 Example 90 Example 91 Example 92 Example 93 Example 95 Example 96 Example 97 Example 97 Example 100 Example 101 Example 102 Example 106 Example 107 Example 108 Example 109 Example 100 Example 100 Example 101 Example 102 Example 104 Example 105 Example 106 Example 107 Example 108 Example 109 Example 110 Example 111 Example 111	130° C.	60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54 0.48 0.60 0.62 0.66 0.63 0.69 0.51 0.55 0.58 0.51 0.50 0.58 0.56 0.46 0.45 0.59 0.50 0.53 0.51 0.55 0.55 0.55 0.55 0.55 0.55 0.55	A A A A A A A A A A A A A A A A A A A	455055	ple 451 452 453 454 455 456 457 458 459 460 461 462	Example 151 Example 152 Example 153 Example 154 Example 155 Example 156 Example 157 Example 157 Example 158 Example 159 Example 160 Example 161 Example 161 Example 161	Heating Temper- ature 130° C.	Time 60 Minutes	of uniformity of coat 0.57 0.53 0.53 0.46 0.52 0.57 0.54 0.56 0.46 0.64 0.64 0.64 0.62	A A A A A A B A A A A
384 385 386 387 388 389 390 391 392 393 394 395 396 397 398 399 400 401 402 403 404 405 406 407 408 409 410 411 411 412 413 414	Example 84 Example 85 Example 86 Example 87 Example 89 Example 90 Example 91 Example 92 Example 93 Example 94 Example 95 Example 96 Example 97 Example 98 Example 100 Example 101 Example 102 Example 105 Example 106 Example 107 Example 106 Example 107 Example 107 Example 108 Example 109 Example 101 Example 101 Example 105 Example 106 Example 107 Example 108 Example 109 Example 110 Example 111 Example 111 Example 111 Example 111	130° C.	60 Minutes	0.59 0.52 0.46 0.51 0.58 0.59 0.56 0.54 0.48 0.60 0.62 0.66 0.63 0.69 0.51 0.55 0.58 0.56 0.46 0.45 0.59 0.50 0.53 0.51 0.55 0.53 0.51 0.55 0.52 0.56 0.60 0.60	A A A A A A A A A A A A A A A A A A A	455055	9le 451 452 453 454 455 456 457 458 459 460 461	Example 151 Example 152 Example 153 Example 154 Example 155 Example 156 Example 157 Example 158 Example 159 Example 160 Example 161 Example 161 Example	Heating Temper- ature 130° C.	Time 60 Minutes	of uniformity of coat 0.57 0.53 0.53 0.46 0.52 0.57 0.54 0.56 0.46 0.46 0.64	A A A A A B A A
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TABLE 18-continued

TABLE 18-continued

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		Heating	condition	Evaluation of					Heating	g condition	Evaluation of	
Exam- ple	Emulsion	Temper- ature	Time	uniformity of coat	Evaluation of image	5	Exam- ple	Emulsion	Temper- ature	Time	uniformity of coat	Evaluation of image
465	Example	130° C.	60 Minutes	0.68	A		502	Example	130° C.	60 Minutes	0.49	A
466	165 Example	130° C.	60 Minutes	0.58	A		503	202 Example	130° C.	60 Minutes	0.54	A
467	166 Example	130° C.	60 Minutes	0.50	В	10	504	203 Example	130° C.	60 Minutes	0.49	A
468	167 Example	130° C.	60 Minutes	0.60	A		505	204 Example	130° C.	60 Minutes	0.55	A
469	168 Example	130° C.	60 Minutes	0.55	В		506	205 Example	130° C.	60 Minutes	0.58	A
470	169 Example	130° C.	60 Minutes	0.48	A	15	507	206 Example	130° C.	60 Minutes	0.58	A
471	170 Example	130° C.	60 Minutes	0.58	A		508	207 Example	130° C.	60 Minutes	0.60	A
472	171 Example	130° C.	60 Minutes	0.48	A		509	208 Example	130° C.	60 Minutes	0.54	A
473	172 Example	130° C.	60 Minutes	0.52	A	20	510	209 Example	130° C.	60 Minutes	0.53	A
474	173 Example	130° C.	60 Minutes	0.48	A		511	210 Example	130° C.	60 Minutes	0.49	A
475	174 Example	130° C.	60 Minutes	0.52	A		512	211 Example	130° C.	60 Minutes	0.60	A
476	175 Example	130° C.	60 Minutes	0.49	A	25	513	212 Example	130° C.	60 Minutes	0.58	A
477	176		60 Minutes			23		213				
	Example 177	130° C.		0.60	A		514	Example 214	130° C.	60 Minutes	0.57	A
478	Example 178	130° C.	60 Minutes	0.45	A .		515	Example 215	130° C.	60 Minutes	0.52	A .
479	Example 179	130° C.	60 Minutes	0.49	A	30	516	Example 216	130° C.	60 Minutes	0.51	A
480	Example 180	130° C.	60 Minutes	0.56	A		517	Example 217	130° C.	60 Minutes	0.47	A
481	Example 181	130° C.	60 Minutes	0.52	A		518	Example 218	130° C.	60 Minutes	0.55	A
482	Example 182	130° C.	60 Minutes	0.52	A	35	519	Example 219	130° C.	60 Minutes	0.67	В
483	Example 183	130° C.	60 Minutes	0.49	A		520	Example 220	130° C.	60 Minutes	0.65	A
484	Example 184	130° C.	60 Minutes	0.52	A		521	Example 221	130° C.	60 Minutes	0.60	Α
485	Example 185	130° C.	60 Minutes	0.54	A	40	522	Example 222	130° C.	60 Minutes	0.66	В
486	Example 186	130° C.	60 Minutes	0.57	A		523	Example 223	130° C.	60 Minutes	0.64	В
487	Example 187	130° C.	60 Minutes	0.51	A		524	Example 224	130° C.	60 Minutes	0.45	В
488	Example 188	130° C.	60 Minutes	0.53	A	45	525	Example 225	130° C.	60 Minutes	0.47	A
489	Example 189	130° C.	60 Minutes	0.66	В		526 527	Example 226 Example 227	130° C. 130° C.	60 Minutes 60 Minutes	0.60 0.46	В А
490	Example	130° C.	60 Minutes	0.69	A		528	Example 228	130° C.	60 Minutes	0.49	В
491	190 Example	130° C.	60 Minutes	0.62	A		529 530	Example 229 Example 230	130° C. 130° C.	60 Minutes 60 Minutes	0.54 0.54	A A
492	191 Example	130° C.	60 Minutes	0.67	В	50	531 532	Example 231 Example 232	130° C. 130° C.	60 Minutes 60 Minutes	0.51 0.51	A A
493	192 Example	130° C.	60 Minutes	0.69	В		533 534	Example 233 Example 234	130° C. 130° C.	60 Minutes 60 Minutes	0.47 0.59	A A
494	193 Example	130° C.	60 Minutes	0.60	A		535 536	Example 235 Example 236	130° C. 130° C.	60 Minutes 60 Minutes	0.51 0.53	A A
495	194 Example	130° C.	60 Minutes	0.66	В	55	537 538	Example 237 Example 238	130° C. 130° C.	60 Minutes 60 Minutes	0.51 0.48	A A
496	195 Example	130° C.	60 Minutes	0.54	A		539 540	Example 239 Example 240	130° C. 130° C.	60 Minutes 60 Minutes	0.57 0.47	A A
497	196 Example	130° C.	60 Minutes	0.49	В		541 542	Example 241 Example 242	130° C. 130° C.	60 Minutes 60 Minutes	0.55 0.54	A A
498	197 Example	130° C.	60 Minutes	0.49	A	60	543 544	Example 243 Example 244	130° C. 130° C. 130° C.	60 Minutes 60 Minutes	0.54 0.54	A A A
	198						545	Example 245	130° C.	60 Minutes	0.47	A
499	Example 199	130° C.	60 Minutes	0.48	В		546 547	Example 246 Example 247	130° C. 130° C.	60 Minutes 60 Minutes	0.53 0.55	A A
500	Example 200	130° C.	60 Minutes	0.50	A		548 549	Example 248 Example 249	130° C. 130° C.	60 Minutes 60 Minutes	0.68 0.57	В А
501	Example 201	130° C.	60 Minutes	0.53	A	65	550 551	Example 250 Example 251	130° C. 130° C.	60 Minutes 60 Minutes	0.47 0.57	В А

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TABLE 18-continued

December 252 130° C. 60 Minutes 0.51 B			Heating	Evaluation of		
553 Example 253 130° C. 60 Minutes 0.58 A 554 Example 254 130° C. 60 Minutes 0.54 A 555 Example 255 130° C. 60 Minutes 0.47 A 556 Example 256 130° C. 60 Minutes 0.48 A 557 Example 257 130° C. 60 Minutes 0.48 A 557 Example 258 130° C. 60 Minutes 0.48 A 559 Example 259 130° C. 60 Minutes 0.47 A 560 Example 261 130° C. 60 Minutes 0.57 A 561 Example 261 130° C. 60 Minutes 0.57 A 562 Example 261 130° C. 60 Minutes 0.59 A 563 Example 263 130° C. 60 Minutes 0.59 A 564 Example 264 130° C. 60 Minutes 0.53 A 565 Example 266 130° C.		Emulsion		Time		Evaluation of image
553 Example 253 130° C. 60 Minutes 0.58 A 554 Example 254 130° C. 60 Minutes 0.54 A 555 Example 255 130° C. 60 Minutes 0.47 A 556 Example 256 130° C. 60 Minutes 0.48 A 557 Example 257 130° C. 60 Minutes 0.48 A 559 Example 259 130° C. 60 Minutes 0.47 A 560 Example 260 130° C. 60 Minutes 0.47 A 561 Example 261 130° C. 60 Minutes 0.47 A 561 Example 261 130° C. 60 Minutes 0.57 A 562 Example 261 130° C. 60 Minutes 0.53 A 563 Example 263 130° C. 60 Minutes 0.59 A 564 Example 264 130° C. 60 Minutes 0.53 A 565 Example 266 130° C.	552	Example 252	130° C.	60 Minutes	0.51	В
554 Example 254 130° C. 60 Minutes 0.54 A 555 Example 255 130° C. 60 Minutes 0.47 A 556 Example 256 130° C. 60 Minutes 0.48 A 557 Example 257 130° C. 60 Minutes 0.48 A 557 Example 258 130° C. 60 Minutes 0.47 A 559 Example 259 130° C. 60 Minutes 0.47 A 560 Example 261 130° C. 60 Minutes 0.57 A 561 Example 261 130° C. 60 Minutes 0.59 A 562 Example 263 130° C. 60 Minutes 0.59 A 563 Example 263 130° C. 60 Minutes 0.59 A 564 Example 264 130° C. 60 Minutes 0.59 A 565 Example 265 130° C. 60 Minutes 0.56 A 566 Example 267 130° C.	553		130° C.	60 Minutes	0.58	A
556 Example 256 130° C. 60 Minutes 0.48 A 557 Example 257 130° C. 60 Minutes 0.56 A 558 Example 258 130° C. 60 Minutes 0.48 A 559 Example 259 130° C. 60 Minutes 0.47 A 560 Example 260 130° C. 60 Minutes 0.57 A 561 Example 261 130° C. 60 Minutes 0.59 A 562 Example 261 130° C. 60 Minutes 0.59 A 563 Example 263 130° C. 60 Minutes 0.59 A 564 Example 264 130° C. 60 Minutes 0.59 A 565 Example 264 130° C. 60 Minutes 0.59 A 565 Example 265 130° C. 60 Minutes 0.53 A 566 Example 266 130° C. 60 Minutes 0.56 A 567 Example 268 130° C.	554		130° C.	60 Minutes	0.54	A
557 Example 257 130° C. 60 Minutes 0.56 A 558 Example 258 130° C. 60 Minutes 0.48 A 559 Example 259 130° C. 60 Minutes 0.47 A 560 Example 260 130° C. 60 Minutes 0.57 A 561 Example 261 130° C. 60 Minutes 0.59 A 562 Example 263 130° C. 60 Minutes 0.53 A 563 Example 263 130° C. 60 Minutes 0.59 A 564 Example 265 130° C. 60 Minutes 0.59 A 565 Example 265 130° C. 60 Minutes 0.59 A 566 Example 265 130° C. 60 Minutes 0.58 A 567 Example 266 130° C. 60 Minutes 0.56 A 568 Example 268 130° C. 60 Minutes 0.57 A 569 Example 269 130° C.	555	Example 255	130° C.	60 Minutes	0.47	A
558 Example 258 130° C. 60 Minutes 0.48 A 559 Example 259 130° C. 60 Minutes 0.47 A 560 Example 261 130° C. 60 Minutes 0.57 A 561 Example 261 130° C. 60 Minutes 0.59 A 562 Example 262 130° C. 60 Minutes 0.59 A 563 Example 263 130° C. 60 Minutes 0.59 A 564 Example 264 130° C. 60 Minutes 0.53 A 565 Example 265 130° C. 60 Minutes 0.53 A 566 Example 266 130° C. 60 Minutes 0.58 A 567 Example 266 130° C. 60 Minutes 0.54 A 568 Example 269 130° C. 60 Minutes 0.57 A 569 Example 270 130° C. 60 Minutes 0.57 A 570 Example 271 130° C.		Example 256		60 Minutes		A
559 Example 259 130° C. 60 Minutes 0.47 A 560 Example 260 130° C. 60 Minutes 0.57 A 561 Example 261 130° C. 60 Minutes 0.59 A 562 Example 261 130° C. 60 Minutes 0.59 A 563 Example 263 130° C. 60 Minutes 0.59 A 564 Example 264 130° C. 60 Minutes 0.53 A 565 Example 265 130° C. 60 Minutes 0.58 A 566 Example 266 130° C. 60 Minutes 0.56 A 567 Example 268 130° C. 60 Minutes 0.54 A 568 Example 269 130° C. 60 Minutes 0.57 A 569 Example 270 130° C. 60 Minutes 0.57 A 570 Example 271 130° C. 60 Minutes 0.57 A 571 Example 271 130° C.		Example 257				A
560 Example 260 130° C. 60 Minutes 0.57 A 561 Example 261 130° C. 60 Minutes 0.59 A 562 Example 262 130° C. 60 Minutes 0.53 A 563 Example 264 130° C. 60 Minutes 0.53 A 564 Example 264 130° C. 60 Minutes 0.58 A 565 Example 265 130° C. 60 Minutes 0.56 A 566 Example 267 130° C. 60 Minutes 0.56 A 567 Example 268 130° C. 60 Minutes 0.56 A 568 Example 268 130° C. 60 Minutes 0.57 A 569 Example 269 130° C. 60 Minutes 0.57 A 570 Example 270 130° C. 60 Minutes 0.57 A 571 Example 271 130° C. 60 Minutes 0.66 B 572 Example 273 130° C.						
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505 P 1 205 1200 C 60 M 1	584	Example 284	130° C.	60 Minutes	0.49	A
585 Example 285 130° C. 60 Minutes 0.53 A	585	Example 285	130° C.	60 Minutes	0.53	A
586 Example 286 130° C. 60 Minutes 0.58 A	586	Example 286	130° C.	60 Minutes	0.58	A
587 Example 287 130° C. 60 Minutes 0.49 A	587	Example 287	130° C.	60 Minutes	0.49	A
588 Example 288 130° C. 60 Minutes 0.57 A	588	Example 288	130° C.	60 Minutes	0.57	A
589 Example 289 130° C. 60 Minutes 0.51 A	589	Example 289	130° C.	60 Minutes	0.51	A
590 Example 290 130° C. 60 Minutes 0.56 A	590	Example 290	130° C.	60 Minutes	0.56	A
591 Example 291 130° C. 60 Minutes 0.54 A	591	Example 291	130° C.	60 Minutes	0.54	A
592 Example 292 130° C. 60 Minutes 0.50 A	592	Example 292	130° C.	60 Minutes	0.50	A
593 Example 293 130° C. 60 Minutes 0.59 A	593	Example 293	130° C.	60 Minutes	0.59	A
594 Example 294 130° C. 60 Minutes 0.56 A	594	Example 294	130° C.	60 Minutes	0.56	A
595 Example 295 130° C. 60 Minutes 0.59 A	595	Example 295	130° C.	60 Minutes	0.59	A
596 Example 296 130° C. 60 Minutes 0.67 B	596	Example 296	130° C.	60 Minutes	0.67	В
597 Example 297 130° C. 60 Minutes 0.45 A	597	Example 297	130° C.	60 Minutes		A
598 Example 298 130° C. 60 Minutes 0.46 A	598		130° C.	60 Minutes	0.46	A
599 Example 299 130° C. 60 Minutes 0.46 A	599	Example 299	130° C.	60 Minutes	0.46	A
600 Example 300 130° C. 60 Minutes 0.45 A	600	Example 300	130° C.	60 Minutes	0.45	A

TABLE 19

60 Minutes

0.58

130° C.

801

Example 701

Compar- ative		Heating	condition	Evalua- tion of unifor-	Evalua-
Exam- ple	Emulsion	Temper- ature	Time	mity of coat	tion of image
7	Comparative Example 1	130° C.	60 Minutes	0.78 μm	D
8	Comparative Example 2	130° C.	60 Minutes	0.72 μm	С
9	Comparative Example 3	130° C.	60 Minutes	0.71 μm	D
10	Comparative Example 4	130° C.	60 Minutes	0.75 μm	D
11	Comparative Example 5	130° C.	60 Minutes	0.78 μm	С

74 TABLE 19-continued

5	Compar- ative		Heating	condition	Evalua- tion of unifor-	Evalua-
	Exam- ple	Emulsion	Temper- ature	Time	mity of coat	tion of image
	12	Comparative Example 6	130° C.	60 Minutes	0.81 μm	D
10	13	Comparative Example 1	130° C.	60 Minutes	0.74 μm	С
	14	Comparative Example 2	130° C.	60 Minutes	0.76 μm	С

By comparison of Examples 301 to 600 with Comparative Examples 7 to 12, in the emulsion having the configuration described in Japanese Patent Application Laid-Open No. 2011-128213, the charge transporting layer formed using the 20 emulsion after leaving for a long time has inferior uniformity of the coat to that of the emulsion according to the present invention prepared using the solution containing the charge transporting substance and the compound that reduces the surface energy, and water. It is thought that coalescence of the oil droplets in the emulsion after longterm preservation causes aggregation of the oil droplets to reduce the uniformity of the oil droplets in the emulsion; thereby, the uniformity of the coat surface after formation of ³⁰ the charge transporting layer is reduced.

Moreover, by comparison of Comparative Examples with Examples 13 and 14, it turns out that compared to the emulsion according to the present invention prepared using 35 the solution containing the charge transporting substance and the compound that reduces the surface energy, and water, the emulsion having the configuration described in Japanese Patent Application Laid-Open No. 2011-128213 may not obtain sufficient uniformity of the coat even if the emulsion is not preserved for a long time. This shows that in the case where the compound that reduces the surface energy is not used, the particle diameter of the emulsion particle is not sufficiently reduced depending on the condition, and it is difficult to obtain sufficient uniformity of the coat after formation of the charge transporting layer.

The image was evaluated as Rank A or B if the surface roughness was less than 0.7 µm in evaluation of uniformity of the coat surface, and the image was evaluated as Rank C or D if the surface roughness was 0.7 µm or more in evaluation of uniformity of the coat surface. Namely, the uniformity of the coat surface corresponds to unevenness of 55 the image.

While the present invention has been described with reference to exemplary embodiments, it is to be understood that the invention is not limited to the disclosed exemplary embodiments. The scope of the following claims is to be accorded the broadest interpretation so as to encompass all such modifications and equivalent structures and functions.

This application claims the benefit of Japanese Patent 65 Application Nos. 2012-058904, filed Mar. 15, 2012, and 2013-039646, filed Feb. 28, 2013, which are hereby incorporated by reference herein in their entirety.

The invention claimed is:

1. A method of producing an electrophotographic photosensitive member which comprises a support and a charge transporting layer formed thereon, comprising the steps of: preparing a solution comprising a charge transporting material; and at least one compound selected from the group consisting of a fluorine-atom-containing polyacrylate, a fluorine-atom-containing polymethacrylate, a polycarbonate having siloxane bond, a polyester lovation of the product of the product

in which R¹¹ represents a hydrogen or a methyl group, R¹² represents an alkylene group, R¹³ represents a perfluoroalkyl group having carbon atoms 4 to 6.

3. A method of producing the electrophotographic photosensitive member according to claim 1, wherein the polycarbonate having siloxane bond is a polycarbonate A comprising a repeating structural unit represented by the following formula (2-1) and a repeating structural unit represented by the following formula (2-3), or a polycarbonate B comprising a repeating structural unit represented by the following formula (2-2) and a repeating structural unit represented by the following formula (2-3).

$$(2-1)$$

$$CH_{2} \xrightarrow{)_{3}} \left(\begin{array}{c} \mathbb{R}^{14} \\ \mathbb{N}^{15} \\ \mathbb{N}^{15} \end{array} \right) \xrightarrow{\mathbb{N}^{16}} \left(CH_{2} \xrightarrow{)_{3}} \right)$$

$$(2-2)$$

$$\begin{array}{c|c}
 & R^{30} & R^{32} & O \\
 & R^{31} & R^{33}
\end{array}$$

50

bond, a silicone oil, a polyolefin, an aliphatic acid, an 45 aliphatic acid amide, and an aliphatic acid ester;

dispersing the solution in water to prepare an emulsion; forming a coat for the charge transporting layer by using the emulsion; and

heating the coat to form the charge transporting layer.

2. A method of producing the electrophotographic photosensitive member according to claim 1, wherein the fluorine-atom-containing polyacrylate and the fluorine-atom-containing polymethacrylate are represented by the following formula (1),

$$\begin{array}{c|c}
R^{11} & & & (1) & 60 \\
\hline
 & & & & \\
\hline
 & & & & \\
R^{13} - R^{12} - O - C & & \\
 & & & & \\
O & & & & \\
\end{array}$$

in which R^{14} to R^{17} each independently represents a methyl group or a phenyl group, m^1 represents number of repetitions of a structure enclosed in brackets, and an average of m^1 in the polycarbonate A ranges from 20 to 100:

R¹⁸ to R²⁹ each independently represents a methyl group or a phenyl group, m², m³, m⁴ and m⁵ each independently represents number of repetitions of a structure enclosed in brackets, an average of m²+m³+m⁴+m⁵ in the polycarbonate B ranges from 0 to 450, Z¹ and Z² each independently represents an ethylene group or a propylene group, and Z³ represents an oxygen atom, an ethylene group or a propylene group; and

X¹ represents a single bond, a methylene group, an ethylidene group, a propylidene group, a phenylethylidene group, a cyclohexylidene group, or an oxygen atom, and R³0 to R³3 each independently represents a hydrogen atom or a methyl group.

4. A method of producing the electrophotographic photosensitive member according to claim 1, wherein the polyester having siloxane bond is a polyester C comprising a repeating structural unit represented by the following formula (3-1) and a repeating structural unit represented by the following formula (3-2),

in which R³⁴ to R³⁷ each independently represents a methyl group or a phenyl group, Y¹ represents a metaphenylene group, a para-phenylene group, or a bivalent group having two para-phenylene groups bonded with an oxygen atom, m⁶ represents number of repetitions of a structure enclosed in brackets, and an average of m⁶ 25 in the polyester C ranges from 20 to 100; and

R³⁸ to R⁴¹ each independently represents a hydrogen atom or a methyl group, X² represents a single bond, a methylene group, an ethylidene group, a propylidene group, a cyclohexylidene group, or an oxygen atom, and Y² represents a meta-phenylene group, para-phenylene group, or a bivalent group having two paraphenylene groups bonded with an oxygen atom.

5. A method of producing the electrophotographic photosensitive member according to claim 1, wherein the polystyrene having siloxane bond is a polystyrene D comprising a repeating structural unit represented by the following formula (4-1) and a repeating structural unit represented by the following formula (4-2),

$$\begin{array}{c} \begin{array}{c} \begin{array}{c} CH_{3} \\ \end{array} \\ \begin{array}{c} CH_{2} \\ \end{array} \\ \begin{array}{c} C\\ \end{array} \\ \begin{array}{c} C$$

in which m⁷ represents an integer selected from 1 to 10, 60 and m⁸ represents an integer selected from 20 to 100.

6. A method of producing the electrophotographic photosensitive member according to claim **1**, wherein the silicone oil is represented by the following formula (5),

in which R⁴² to R⁴⁵ each independently represents a methyl group or a phenyl group, and m⁹ is an integer selected from 20 to 100.

7. A method of producing the electrophotographic photosensitive member according to claim 1,

wherein the polyolefin is an aliphatic hydrocarbon having carbon atoms 10 to 40.

8. A method of producing the electrophotographic photosensitive member according to claim **1**, wherein the aliphatic acid, the aliphatic acid amide and the aliphatic acid ester are represented by the following formula (7-1),

$$\begin{array}{c} O \\ \parallel \\ \mathbb{R}^{46} - \mathbb{C} - O - \mathbb{R}^{47} \end{array}$$

in which R^{46} represents an alkyl group having carbon atoms 10 to 40, and R^{47} represents a hydrogen atom, an amino group, or an alkyl group having carbon atoms 10 to 40.

9. A method of producing the electrophotographic photosensitive member according to claim **1**,

wherein, in the emulsion, the ratio of a mass of water to a mass of the solution is 5/5 to 7/3.

10. A method of producing the electrophotographic photosensitive member according claim 1, wherein the solution further comprises a binder resin, the binder resin being a polycarbonate resin free from a siloxane bond or a polyester resin free from a siloxane bond.

11. A method of producing the electrophotographic photosensitive member according to claim 1,

wherein the solution further comprises a liquid whose solubility in water under 25° C. and 1 atmosphere is 1.0 mass % or less.

12. An emulsion for a charge transporting layer in which a solution is dispersed in water, wherein the solution comprises:

a charge transporting material; and

at least one compound selected from the group consisting of a fluorine-atom-containing polyacrylate, a fluorine-atom-containing polymethacrylate, a polycarbonate having siloxane bond, a polyester having siloxane bond, a polystyrene having siloxane bond, a silicone oil, a polyolefin, an aliphatic acid, an aliphatic acid amide and an aliphatic acid ester.

 ${f 13}.$ The emulsion for a charge transporting layer according to claim ${f 12},$

wherein the solution further comprises a liquid whose solubility in water under 25° C. and 1 atmosphere is 1.0 mass % or less.

* * * * *